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Quantum search with hybrid adiabatic–quantum-walk algorithms and realistic noise

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Computing using a continuous-time evolution, based on the natural interaction Hamiltonian of the quantum computer hardware, is a promising route to building useful quantum computers in the near term. Adiabatic quantum computing, quantum annealing, computation by a continuous-time quantum walk, and special purpose quantum simulators all use this strategy. In this work, we carry out a detailed examination of adiabatic and quantum-walk implementation of the quantum search algorithm, using the more physically realistic hypercube connectivity, rather than the complete graph, for our base Hamiltonian. We calculate optimal adiabatic schedules both analytically and numerically for the hypercube and then interpolate between adiabatic and quantum-walk searching, obtaining a family of hybrid algorithms. We show that all of these hybrid algorithms provide the quadratic quantum speedup when run with optimal parameter settings, which we determine and discuss in detail. We incorporate the effects of multiple runs of the same algorithm, noise applied to the qubits, and two types of problem misspecification, determining the optimal hybrid algorithm for each case. Our results reveal a rich structure of how these different computational mechanisms operate and should be balanced in different scenarios. For large systems with low noise and good control, a quantum walk is the best choice, while hybrid strategies can mitigate the effects of many shortcomings in hardware and problem misspecification.

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I. INTRODUCTION

Quantum computing based not on discrete quantum gates, but on continuous-time evolution under quantum Hamiltonians, is a promising route towards near-future useful quantum computers. This is in part because of the success of experimental quantum annealing efforts [1–5], and also because of special purpose quantum simulators [6,7] that employ this technique, and are potentially useful for a wider range of computations. Problems known to be suitable for continuous-time algorithms are wide ranging across many important areas, including finance [8], aerospace [9], machine learning [10–12], theoretical computer science [13], decoding of communications [14], mathematics [15,16], and computational biology [17].

Continuous-time computation is less familiar than the ubiquitous digital computation that underpins everything from mobile phones to internet servers. There is no classical equivalent of computing via continuous-time manipulation of digital data to guide our intuition or provide a source of classical algorithmic resources that might be adapted to a quantum setting. A detailed study based on a well-characterized problem can thus serve to elucidate the mechanisms in continuous-time quantum computing and build a firm foundation for further development. Hence, we focus this work on the unordered search problem, which was first studied in the quantum setting by Grover [18]. Grover’s algorithm provides a quadratic speedup over classical searching, proved by Bennett et al. [19] to be the best possible improvement.

Two further examples of quantum search algorithms are quantum-walk (QW) searching [20] and the adiabatic quantum computing (AQC) search algorithm [21], which both obtain the optimal quadratic speedup. There remain the questions of which is more efficient in terms of the prefactors [22] or more robust in the face of imperfections. While the results in [19] imply that any protocol we develop here will not provide better scaling properties, asymptotic scaling factors do not give a full account of algorithm performance. A recent study in which a quantum annealer appears to show the same asymptotic scaling as a classical algorithm, but with a prefactor advantage [3] of \(10^8\), underscores the importance of practical computational advantages beyond asymptotic scaling. This prompts more detailed study of exactly how the quantum search algorithms work, the topic of many papers since the original algorithms were presented [23–26].

Quantum-walk searching has been shown to implement a similar type of rotation in Hilbert space to that which Grover’s algorithm employs [20]. On the other hand, adiabatic quantum searching alters the Hamiltonian over time, turning on the term for the marked state slowly enough to keep the quantum system in its ground state throughout. On the face of it, these are quite different dynamics, as has been highlighted in [27]. However, both use the same Hamiltonians and initial states and we argue here that both are best viewed as extreme cases of possible quantum annealing schedules. This invites...
consideration of intermediate quantum annealing schedules, and we show how to interpolate smoothly between QW and AQC searching, enabling both mechanisms to contribute to solving the search problem. We examine the hybrid algorithms thus created using simplified models for the asymptotic scaling, and numerical simulation to explore smaller systems where more complex finite-size effects contribute. Taking into account realistic factors, such as a finite initialization time for each run of the algorithm, our results reveal a rich structure of intermediate strategies available to optimize the performance of a practical quantum computer.

The paper is structured as follows. In Sec. II we give the background and lay the groundwork for our study in terms of the QW and AQC protocols which we interpolate between. In Sec. III we introduce the two AQC schedules which we use in this study and we explain in detail how they arise from the dynamics of the quantum search Hamiltonian on a hypercube. In Sec. IV we construct interpolated protocols which can take advantage of both QW and AQC mechanisms. We then turn to the performance of the interpolated protocols in finite-size systems. In Sec. IV C we examine the scaling for larger systems in detail and demonstrate that the interpolated protocols also yield a quadratic speedup over classical searching. In Sec. V we incorporate strategies which involve performing multiple runs, including in Sec. V C the effect of adding decoherence, and in Sec. VI we examine the effect of problem mis specification. Finally, in Sec. VII we summarize our results and their implications for future work. The calculation of the optimal schedule for the hypercube is outlined in Appendix A and notes on our numerical methods are in Appendix B.

II. BACKGROUND

We begin with a discussion of how the unstructured search problem may be encoded into qubit states. From here we show how, with the use of very similar Hamiltonians, the search problem can be solved with an optimal quantum scaling advantage, via both QW and AQC algorithms.

A. Encoding search into quantum states

The search problem can be framed in terms of the $N = 2^n$ basis states of an $n$-qubit system $|j\rangle = |\{0\}, |1\rangle |m\rangle$, where $|\{0\}, |1\rangle$ is the basis of a single qubit. We are given that one of the basis states behaves differently from the others and denote this “marked” state by $|m\rangle$, where $m$ is an $n$-digit bit string identifying one of the basis states. Because of the difference in behavior, we can easily verify whether a given state is the marked state. One way to implement this is for the marked state to have a lower energy than all other states, e.g., using a Hamiltonian like $\hat{H}_G = \mathbb{1} − |m\rangle \langle m|$, where $\mathbb{1}$ is the identity operator. In terms of Pauli operators, $\hat{H}_G = \mathbb{1} − \frac{1}{2^n} \sum_{j=1}^{n} (q_j \sigma_j^z + \mathbb{1})$,  

where $q_j \in \{-1, 1\}$ define a logical bit string $m$ via the mapping $1 \to 0$ and $-1 \to 1$. The search problem is then to determine which of the basis labels $j$ corresponds to the marked state label $m$, given that a priori we have no knowledge of $m$, apart from it being a basis state. We represent this ignorance of the marked state by starting with the system in a uniform superposition over the basis states, $|\psi_{\text{init}}\rangle = \frac{1}{\sqrt{N}} \sum_{j=0}^{N-1} |j\rangle$.

The quantum search algorithms considered in this paper solve the search problem by evolving the system into a state with a large overlap with the marked state so that a measurement can be made to return the marked state label $m$ with high probability. This is achieved by applying a (generally time-dependent) Hamiltonian to evolve the system initially in state $|\psi_{\text{init}}\rangle$ to a final state $|\psi_{\text{final}}\rangle$. Performing a measurement of this state in the basis $|\{j\}\rangle$ will yield the marked state label with probability $|\langle \psi_{\text{final}} |m\rangle|^2$. If $|\langle \psi_{\text{final}} |m\rangle|^2 = 1$ then the search is perfect and the problem is solved. If the search is imperfect then the problem can be solved by searching multiple times: Since the result of each search is checked independently, a single successful search is sufficient. As long as $|\langle \psi_{\text{final}} |m\rangle|^2$ is greater than $1/poly(n)$ this form of amplitude amplification will be efficient. Multiple runs have a cost; see Sec. V for details of the tradeoff between multiple runs and the initialization time for each run.

In general, problems with full permutation symmetry, such as the search problem, are considered to be toy problems from a practical point of view. A naive implementation of such a problem, in this case $\hat{H}_G = \mathbb{1} − |m\rangle \langle m|$, the marked state Hamiltonian, requires exponentially many terms of the form $\prod_{j \neq m} \hat{\sigma}_j^{(z)}$, where $m$ is a binary number with $n$ bits (with $j \in m$ indicating the 1 digits of the number) and $j$ iterates over the bits in $m$ that are equal to one. However, it has recently been shown [13] that the spectrum of such terms in permutation-symmetric problems can be reproduced using $n$ extra qubits and a number of extra coupling terms of the form $\hat{\sigma}_j^{(z)} \hat{\sigma}_k^{(z)}$ which scales as $n^2$. It has also been suggested that such models may be fully realized perturbatively [28,29]. Although this approach to construct such terms is much closer to the realm of what can be experimentally realized, it would still be highly nontrivial to implement. Nonetheless, the insights gained from studying the search problem can be adapted to realistic problems of practical interest.

B. Quantum-walk-search algorithm

A continuous-time quantum walk can be defined by considering the labels $j$ of the $n$-qubit basis states $|\{j\}\rangle$ to be the labels of vertices of an undirected graph $G$. The edges of $G$ can be defined through its adjacency matrix $A$, whose elements satisfy $A_{jk} = 1$ if an edge in $G$ connects vertices $j$ and $k$ and $A_{jk} = 0$ otherwise. Since $G$ is undirected, $A$ is symmetric, hence it can be used to define a Hamiltonian. Although we can use the adjacency matrix $A$ directly, it is in general more convenient mathematically to define the Hamiltonian of the quantum walk using the Laplacian $L = A - D$, where $D$ is a diagonal matrix with entries $D_{jj} = d_j$ the degree of vertex $j$ in the graph. We follow this convention here, but note that in this work we use regular graphs for which the degree $d_j = d$ is the same for all vertices, so that $D = d I$, where $I$ is the identity matrix (ones on the diagonal) of the same dimension as $A$. 

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Terms proportional to the identity in the Hamiltonian shift the zero point of the energy scale and contribute an unobservable global phase, but otherwise do not affect the dynamics. The quantum-walk Hamiltonian is then defined as \( \hat{H}_{QW} = -\gamma \hat{L} \), where \( \hat{L} \) is the Laplacian operator and the prefactor \( \gamma \) is the hopping rate of the quantum walk. For any regular graph of degree \( d \) we thus have

\[
\hat{H}_{QW} = \gamma \left( \hat{d} - \sum_{jk} A_{jk} |j\rangle \langle k| \right) \equiv \gamma (\hat{d} - \hat{\Lambda}),
\]

(3)

where the adjacency operator \( \hat{\Lambda} \) has matrix elements in the vertex basis \( |j\rangle \) given by the adjacency matrix \( A \). The action of \( \hat{H}_{QW} \) is to move amplitude between connected vertices, as specified by the nonzero entries in \( A \). During a quantum walk, a pure state \( |\psi(0)\rangle \) evolves according to the Schrödinger equation to give

\[
|\psi(t)\rangle = \exp(-i\hat{H}_{QW}t)|\psi(0)\rangle
\]

(4)

after a time \( t \), where we have used units in which \( \hbar = 1 \).

Quantum-walk dynamics can be used to solve the search problem by modifying the energy of the marked state \( |m\rangle \) to give a quantum-walk-search Hamiltonian

\[
\hat{H}_{QWS} = \gamma (\hat{d} - \hat{\Lambda}) - |m\rangle \langle m|.
\]

(5)

In the units we are using, this amounts to giving state \( |m\rangle \) an energy of \(-1\) while all other states have zero energy. This also makes \( \gamma \) a dimensionless parameter controlling the ratio of the strengths of the two parts of the quantum-walk-search Hamiltonian. Applying \( \hat{H}_{QWS} \) to the search initial state \( |\psi_{\text{init}}\rangle \) in Eq. (2) produces a periodic evolution such that the overlap with the marked state oscillates. The frequency of these oscillations depends on the hopping rate \( \gamma \), which must be chosen correctly, along with the measurement time \( t_f \), to maximize the final success probability \( P = |\langle \psi(t_f)|m\rangle|^2 \), where \( |\psi(t_f)\rangle = \exp(-i\hat{H}_{QWS}t_f)|\psi_{\text{init}}\rangle \) is the state at time \( t_f \).

The performance of quantum-walk-search algorithms will clearly have some dependence on the choice of the graph \( G \). Provided the connectivity is not too sparse or low-dimensional [30], most choices of graph will work, even random graphs [31]. Two convenient choices on which the quantum walk is analytically solvable are the complete graph, for which all vertices are directly connected, and a graph whose edges form an \( n \)-dimensional hypercube. Moore and Russell [32] studied quantum walks on hypercubes and Hein and Tanner [33] performed a detailed analysis of discrete-time quantum-walk searching on the hypercube, extending the work of Shenvi et al. [20]. We choose to focus our work on a hypercube, rather than a fully connected graph, because it is the more practical graph in terms of implementation on a quantum computer. A hypercube graph is the natural choice for a quantum walk encoded into qubits because moving from one vertex to a neighboring vertex corresponds to flipping a qubit. The techniques and scaling arguments we give in this work also apply in the case of a fully connected graph and can be easily extended to a more general setting, for example, to the “typical” random graphs considered in [31].

The adjacency matrix of an \( n \)-dimensional hypercube graph has elements \( A_{jk} = 1 \) if and only if the vertex labels \( j \) and \( k \) have a Hamming distance of one. That is, when written as \( n \)-digit bit strings, they differ in exactly one bit position. The corresponding adjacency operator can be conveniently expressed as

\[
\hat{\Delta}^{(h)} = \sum_{j=1}^{n} \hat{\sigma}_x^{(j)},
\]

(6)

where the sum is over all \( n \) qubits and \( \hat{\sigma}_x^{(j)} \) is the Pauli-X operator applied to the \( j \)th qubit with the identity operator on the other qubits. That is,

\[
\hat{\sigma}_x^{(j)} = \left( \bigotimes_{r=1}^{j-1} \hat{1}_2 \right) \otimes \hat{\sigma}_x \otimes \left( \bigotimes_{r=j+1}^{n} \hat{1}_2 \right),
\]

(7)

where \( \otimes \) denotes the tensor product and \( \hat{1}_2 \) is the identity operator of dimension 2. The Hamiltonian for the quantum walk on the hypercube is thus given by

\[
\hat{H}_{QWS}^{(h)} = \gamma \left( n\hat{d} - \sum_{j=1}^{n} \hat{\sigma}_x^{(j)} \right),
\]

(8)

since an \( n \)-dimensional hypercube has vertices which have degree \( n \).

To construct the quantum-walk-search Hamiltonian on the hypercube, we include two trivial adjustments for later mathematical convenience. If we make the energy of the marked state lower by adding \( -|m\rangle \langle m| \) to the quantum-walk Hamiltonian, this gives the marked state an energy of zero while all other states have an energy of one for this part of the Hamiltonian. Here, we define the hopping rate \( \gamma \) with an explicit factor of \( \frac{1}{\pi} \) to match Refs. [30,34,35] and facilitate the mapping to the symmetric subspace (Appendix A). Our quantum-walk search on the hypercube is then

\[
\hat{H}_{QWS}^{(h)} = \gamma \left( n\hat{d} - \sum_{j=1}^{n} \hat{\sigma}_x^{(j)} \right) + (1 - |m\rangle \langle m|).
\]

(9)

Childs and Goldstone [30] analyze the quantum-walk-search algorithm for both the complete and hypercube graphs. For each graph, they find optimal values of \( \gamma \) for which the performance of the search matches the quadratic quantum speedup achieved by Grover’s search algorithm. The mechanism for finding the marked state can be understood intuitively as follows. Note that the initial state \( |\psi_{\text{init}}\rangle \) from Eq. (2) is the (nondegenerate) ground state of both the complete graph and the hypercube Hamiltonians, i.e., \( \hat{H}_{QWS}^{(h)} \) of Eq. (8). The marked state \( |m\rangle \) is, by design, the ground state of the marked state component of the search Hamiltonian. For large values of \( \gamma \), the marked state term is relatively small, so the graph Hamiltonian dominates, and the ground state of the full search Hamiltonian \( \hat{H}_{QWS}^{(h)} \) of Eq. (9) is approximately \( |\psi_{\text{init}}\rangle \). Conversely, for small values of \( \gamma \), the ground state of \( \hat{H}_{QWS}^{(h)} \) is approximately \( |m\rangle \). Over a narrow range of intermediate values of \( \gamma \), the ground state switches between the two. By calculating the low-level part of the energy spectrum of \( \hat{H}_{QWS}^{(h)} \), Childs and Goldstone tune \( \gamma \) until both the initial state \( |\psi_{\text{init}}\rangle \) and the marked state \( |m\rangle \) have significant overlap with both the ground state \( E_0 \) and the first excited state \( E_1 \) of the search
Hamiltonian. Intuitively, we want to search the Hamiltonian to drive transitions between \(|\psi_{\text{init}}\rangle\) and \(|m\rangle\) as efficiently as possible. This occurs when the overlaps are evenly balanced, which in turn occurs when the gap \(g = E_1 - E_0\) between the ground and first excited state is smallest: \(g_{\text{min}}\). With this optimally chosen value of \(g\), the time it takes for the transition to occur turns out to be proportional to \(1/g_{\text{min}}\). For the hypercube graph, the optimal value of \(g\) is

\[
\gamma_{0}^{(b)} = \frac{1}{N} \sum_{r=0}^{n} \binom{n}{r} \frac{1}{r} \equiv R_1, \tag{10}
\]

where \(\binom{n}{r}\) is the binomial coefficient \(n\) choose \(r\). This sum appears many times in the following calculations, so it is convenient to abbreviate it by \(R_1\). Note also that it is not always sufficiently accurate to use the approximation \(R_1 \approx 2/N\) given in [30]. The time to reach the first maximum overlap with the marked state is \(t_v^{(b)} \approx (\pi/2)\sqrt{N}\), providing a quadratic speedup equivalent to Grover’s original search algorithm.

C. Adiabatic quantum search algorithm

Adiabatic quantum computing, introduced by Farhi et al. [34], works as follows. The problem of interest is encoded into an \(n\)-qubit Hamiltonian \(\hat{H}_p\) in such a way that the solution can be derived from the ground state of \(\hat{H}_p\). The system is initialized in the ground state of a different Hamiltonian \(\hat{H}_0\), for which this initialization is easy. The computation then proceeds by implementing a time-dependent Hamiltonian that is transformed slowly from \(\hat{H}_0\) to \(\hat{H}_p\). In general, this adiabatic “sweep” Hamiltonian can be parametrized in terms of a time-dependent schedule function \(s \in [0, 1]\) as

\[
\hat{H}_{\text{AQQC}}(s) = (1 - s)\hat{H}_0 + s\hat{H}_p, \tag{11}
\]

with \(s = s(t)\) such that \(s(0) = 0\) and at the final time \(t_f\) we have \(s(t = t_f) = 1\). It is useful to define a reduced time \(\tau = t/t_f\), with \(0 \leq \tau \leq 1\). Whereas \(\tau\) is linear in \(t\), the schedule function \(s(\tau)\), written as a function of \(t\) or \(\tau\), allows for nonlinear transformation. Nonlinear schedules are essential to obtain a quantum speedup.

The adiabatic theorem of quantum mechanics [36] says that the system will stay in the instantaneous ground state of the time-dependent Hamiltonian \(\hat{H}_{\text{AQQC}}(s)\) provided the following two conditions are satisfied: (i) There is at all times an energy gap \(g(s) > 0\) between the instantaneous ground and first excited states and (ii) the Hamiltonian is changed sufficiently slowly. Provided these are both true, the system will be in the desired ground state of \(\hat{H}_p\) at the end of the computation, thus solving the problem encoded in \(\hat{H}_p\). In practice, the duration of this adiabatic sweep would be prohibitively long, so a realistic sweep will incur some probability of error. We discuss this and other subtleties of the adiabatic theorem in Sec. III, after we introduce the adiabatic quantum search algorithm. For a comprehensive overview of AQQC, see Ref. [37].

Roland and Cerf [21] describe how adiabatic quantum computing can be used to solve the search problem with a quadratic quantum speedup. Define the problem Hamiltonian as

\[
\hat{H}_0 = \mathbb{1} - |m\rangle\langle m|, \tag{12}
\]

whose nondegenerate ground state is equal to the marked state \(|m\rangle\) with eigenvalue zero. We then need to choose our easy Hamiltonian \(\hat{H}_0\) such that it has \(|\psi_{\text{init}}\rangle\) as defined in Eq. (2), as its nondegenerate ground state. There are many possible choices; Roland and Cerf use \(\hat{H}_0 = \mathbb{1} - |\psi_{\text{init}}\rangle\langle \psi_{\text{init}}|\). With the system initialized in \(|\psi_{\text{init}}\rangle\), the algorithm proceeds by implementing the time-dependent Hamiltonian in Eq. (11), with a suitable schedule function \(s(\tau)\), so that after a time \(t_f\) the final state of the system is close to the marked state \(|m\rangle\). Roland and Cerf demonstrate that a linear schedule function \(s(\tau) = \tau = t/t_f\) does not produce a quantum speedup. It is necessary to use a more efficient nonlinear \(s(\tau)\), whose rate of change is in proportion to the size of the gap \(g(s)\) at that point in the schedule, in order to produce the quadratic speedup of Grover’s search algorithm.

It is easy to show that \(\hat{H}_0 = \mathbb{1} - |\psi_{\text{init}}\rangle\langle \psi_{\text{init}}|\) is proportional to the adjacency operator of the fully connected graph with \(N = 2^n\) vertices. For the reasons already given in the context of the quantum-walk-search algorithm, a Hamiltonian corresponding to a less connected graph is preferable for practical applications. In order to make direct comparisons between adiabatic and quantum-walk searching, we use the hypercube graph, since this also has \(|\psi_{\text{init}}\rangle\) as its nondegenerate ground state, with Hamiltonian (in its Laplacian form) given by

\[
\hat{H}_0^{(b)} = \frac{1}{2} \left( n\mathbb{1} - \sum_{j=1}^{n} \hat{\sigma}_j^{(j)} \right), \tag{13}
\]

where we have again included a factor of one-half for mathematical convenience. As further motivation for this choice, we note that this corresponds to a transverse-field driver Hamiltonian applied to qubits, which is the most common choice for quantum annealing hardware and which can be experimentally realized on a large scale [38]. Combining Eqs. (12) and (13), we have the adiabatic quantum computing Hamiltonian for search on a hypercube,

\[
\hat{H}_{\text{AQQC}}^{(b)} = (1 - s) \frac{1}{2} \left( n\mathbb{1} - \sum_{j=1}^{n} \hat{\sigma}_j^{(j)} \right) + s(1 - |m\rangle\langle m|). \tag{14}
\]

We note that \(\hat{H}_{\text{AQQC}}^{(b)}\) contains the same terms as \(\hat{H}_{\text{QWS}}^{(b)}\) in Eq. (9), only in different, time-varying proportions. It remains to specify the function \(s(\tau)\) for the optimal performance of this Hamiltonian for searching. There are several subtleties to deriving an optimal \(s(\tau)\) for the hypercube, which we address in the next section.

III. OPTIMIZING AQQC SCHEDULES

We have seen that QW and AQQC searching may be achieved with Hamiltonians that have the same terms but different, time-varying, coefficients. Next we would like to interpolate these coefficients to generate hybrid search algorithms. However, we must first determine an optimal schedule \(s(\tau)\) for the AQQC search. In fact, this is not entirely straightforward: It is possible to find more than one optimal schedule. In this section we derive two different schedules via an analytical method and a numerical method and demonstrate
that these both give optimal quantum scaling advantage for the unstructured search problem.

A. Adiabatic condition and method

We now return to the nuances of the adiabatic theorem and how, in the regime of limited running time, the schedule \( s(\tau) \) may be optimized to minimize the error. A more quantitative statement of the adiabatic theorem [21,34,37,39,40] proceeds as follows: Consider a time-dependent Hamiltonian of the form in Eq. (11), with initial and final Hamiltonians \( \hat{H}_0 \) and \( \hat{H}_f \), respectively, and parametrized by the schedule function \( s(\tau) \) that sweeps from \( s(0) = 0 \) to \( s(1) = 1 \) over a time \( t_f \), the runtime of the sweep. Denote by \( |E_j(t)\rangle \) the \( j \)-th energy eigenstate of the Hamiltonian at time \( t \) and its energy by \( E_j(t) \), where \( j = 0, 1 \) denotes the ground and first excited states, respectively. Provided that \( E_j(t) > E_0(t) \) for \( t \in [0, t_f] \) and transitions to higher energy eigenstates can be ignored, the final state obeys

\[
|\langle \psi(t_f)|E_0(t_f)\rangle|^2 \geq 1 - \epsilon^2
\]

for small parameter \( \epsilon \ll 1 \), provided that at all times

\[
|\frac{d\hat{H}}{dt}\rangle|_{0,1} \leq \epsilon \ll 1,
\]

where the matrix element \( \langle \hat{H}\rangle\rangle|_{0,1} \) is given by

\[
\langle \frac{d\hat{H}}{dt}\rangle|_{0,1} = \left(E_0(t)\left|\frac{d\hat{H}}{dt}\right|E_1(t)\right)
\]

and the gap \( g(t) \) is given by

\[
g(t) = E_1(t) - E_0(t).
\]

However, adiabatic protocols derived from Eq. (16) are not always optimal. This equation accounts for probability amplitude leaking from the ground state into a nearly empty first excited state. Thus it will break down in situations where transitions to higher excited states are important or where the population of the first excited state is significant. We can therefore describe Eq. (16) as a two-level approximation. In the context of the search algorithms studied here, such an approximation turns out to be good for all but the smallest values of \( n \) and becomes more accurate for larger search spaces. We make extensive use of this in what follows, especially in Sec. IV E.

Equation (16) also does not take into account the return of probability amplitude which has already entered the excited state. Such effects can become the most relevant to the dynamics under two circumstances. If the first excited state is populated significantly, then nonadiabatic dynamics can occur such that this amplitude returns and interferes with the ground-state amplitude. This is the regime which we primarily study in this work. Quantum-walk dynamics are an extreme example of such behavior as they can be viewed as time-independent coherent evolution bracketed by instantaneous quenches, which are the ultimate nonadiabatic transitions. The second and more subtle case is deep in the adiabatic regime, where the Hamiltonian sweep rate is so slow that the rate of excitation formation is very low during the middle of the anneal. In these cases, boundary effects become important, which depend in a complicated way on both the nature of the annealing schedule and the total runtime [41–43]. While this regime is very interesting, it is outside the scope of this study and not relevant for practical implementation of algorithms. For this reason, we limit our numerical studies to a maximum runtime of \( \sim 5\pi/\delta_{\text{min}} \), about ten times the typical runtime derived from the minimum gap. With runtimes \( t_f \lesssim 5\pi/\delta_{\text{min}} \), we do not observe any appreciable boundary effects in our numerical results.

Roland and Cerf [21] derived a schedule \( s(\tau) \) for the fully connected graph by optimizing Eq. (16), by matching the instantaneous rate of change of the schedule function \( s(\tau) \) to the size of the gap at that time. Using

\[
\left\langle \frac{d\hat{H}}{dt}\right\rangle_{0,1} = \frac{ds}{dt}\left|\frac{d\hat{H}}{ds}\right|_{0,1},
\]

in the adiabatic condition of Eq. (16) gives

\[
\frac{ds}{dt} \leq \epsilon \frac{g^2(t)}{\delta_{\min}}.
\]

The instantaneous gap \( g(t) \) and \( \langle d\hat{H}/ds\rangle_{0,1} \) can be calculated from the eigensystem of the Hamiltonian, which is analytically tractable for the complete graph. The schedule they obtain this way produces the full quadratic quantum speedup for the adiabatic quantum search algorithm on the fully connected graph.

B. Hypercube schedule calculation

Since we are using the hypercube graph, we must do the equivalent calculation for the hypercube AQC search Hamiltonian given by Eq. (14). The eigensystem of this Hamiltonian has been solved in Refs. [30,34,35] by mapping it to the symmetric subspace. From here the position and size of the minimum gap can be found exactly and the eigenvalue equations expanded about this point. This is combined with the saturation of Eq. (20) at the minimum gap point, where the right-hand side takes its minimum value. From the resulting expressions it is possible to derive an analytical expression for the schedule \( s(\tau) \). The full calculation is somewhat lengthy and is outlined in Appendix A. We find the calculated optimal schedule

\[
s^{(\text{c})}(t) = \frac{2\sqrt{R_2}}{\sqrt{N}(1+R_1)} \tan \left\{ \frac{8\epsilon \sqrt{R_2^2 R_1}}{n\sqrt{R_2^2}} - c \right\} + \frac{1}{1+R_1},
\]

where terms \( O(1/N) \) and smaller have been dropped,

\[
c = \arctan \left( \frac{(1+R_1)\sqrt{N}}{2\sqrt{R_2}} \right),
\]

and the constant \( R_1 \) is defined in Eq. (10) and \( R_2 \) by

\[
R_2 = \frac{1}{N} \sum_{r=1}^{n} \left( \begin{array}{c} n \\ r \end{array} \right) \frac{1}{r^2}.
\]

This analytical schedule is guaranteed to satisfy Eq. (20) only in the region of the minimum gap, however it is here that transitions to unwanted higher energy levels are most rapid,
so the net effect is that this schedule still manages to produce optimal quantum speedup. For \( N \gg 1 \), the runtime is given by

\[
\epsilon_f^{(c)} \simeq \frac{\pi \sqrt{N}}{4},
\]  

where the approximation of the arctangents by \( \pi/2 \) becomes exact as \( N \to \infty \). Note that choosing a value for \( \epsilon \), the accuracy with which the system stays in the ground state [see Eq. (15)], determines the corresponding runtime \( t_f \) and vice versa. For our numerical calculations we have chosen to specify \( t_f \), since this enables direct comparisons with QW searching to be made. The energy levels of \( \hat{H}_{AQC}^{(b)} \) are shown in Fig. 1(a) for \( n = 9 \) and for comparison the energy levels of the search Hamiltonian for the complete graph (which is the same for any size) are shown in Fig. 1(b).

We also solve Eq. (16) numerically to obtain \( s^{(n)} \) using an explicit numerical calculation of the gap \( g(t) \) and using the maximum value of \( (d\hat{H}_{AQC}^{(b)}/dt)_{0,1} \), which is shown in Appendix A to be \( n/4 \). Our numerical algorithm is described in Appendix B. While it does not provide a closed-form solution, results using \( s^{(n)} \) do provide insight into the accuracy of \( s^{(c)} \). Provided the numerics are performed to a sufficient accuracy, \( s^{(n)} \) will always provide an optimal \( \sqrt{N} \) speed up. The analytically and numerically calculated gaps are plotted in Fig. 1(c) for \( n = 9 \) and the corresponding gap for the complete graph is shown in Fig. 1(d). For the hypercube, the analytical and numerical gaps are strikingly different, yet both produce schedules that obtain a quantum speedup. As we will see, this is because for the quantum search problem only the position and size of the gap are important. Elsewhere, the transition probabilities are so small it does not matter how fast the schedule proceeds.

However, note that both of these schedules assume a two-level approximation, as they start from Eq. (16). While in general for large \( N \) this is a good approximation, for small system sizes the higher-energy levels do affect the performance, as we show in the next section.

### C. Performance of hypercube schedules

Having calculated optimal schedules both analytically and numerically, we now compare their performance for system sizes up to \( n = 20 \) qubits. Note that the size of the minimum gap \( g_{(\text{min})} \) calculated from the two-level approximation in Sec. III is exactly the same for both. Since both are based only on the interactions of the two lowest-energy levels, both will find the correct shape for the annealing protocol in this region. Numerical results support this prediction in that for \( n = 20 \) the numerically calculated optimal schedule \( s^{(n)} \) slows down at the same value as \( s^{(c)} \) in Fig. 2. For \( n = 5 \) qubits the schedules are distinct over the whole range of \( \tau \), while for \( n = 20 \) the schedules are almost identical, the only visible difference occurring at \( \tau \lesssim 0.1 \). The difference between them around \( \tau \lesssim 0.1 \) is likely due to interactions with the higher excited states of the hypercube Hamiltonian early in the schedule. In the large-system limit this difference will have little effect on the overall success probability, as the overlap with the initial ground state and the manifold of states participating in the avoided crossing approaches one exponentially fast in the number of qubits \( n \) (see Table I).

For \( n < 20 \), the difference between \( s^{(c)} \) and \( s^{(n)} \) at early times in the run does affect their relative performance, as Fig. 3 shows. Although the numerical schedule \( s^{(n)} \) is a more accurate solution of the optimization in Eq. (16), \( s^{(c)} \) does better than \( s^{(n)} \). The reason is that, while the gap is relatively small early in the schedule, so is the matrix element between the ground and first excited state of the marked state Hamiltonian, as shown in the top inset of Fig. 3. As a result, the numerically calculated schedule slows down unnecessarily in this region, as can be seen in Fig. 2. The approximate expansion for the gap used to derive the schedule \( s^{(c)} \) in Appendix A grows within this region (see Fig. 1). Hence, \( s^{(c)} \) traverses this part of the schedule much faster than \( s^{(n)} \). Effectively, the approximate nature of the expansion for the gap used

![Figure 1](image1.png)

![Figure 2](image2.png)
TABLE I. Numerical fits for various quantities related to quantum walks and adiabatic protocols. These fits were performed using linear fitting on either logarithmic or semilogarithmic axes in the range \( n = 40–70 \), except for \([|\psi_{\text{null}}|E_0|^2 + |\psi_{\text{null}}|E_1|^2]\), which was fit over the range \( n = 11–40 \) due to numerical precision issues. The coefficient of determination \( r^2 = 1 - \frac{\sum_{i=0}^{n-1} y_i - \hat{y}_i}{\sum_{i=0}^{n-1} y_i - \bar{y}} \), where \( f_i \) are the data and \( y \) is the fitting function, is calculated against the linear function on the logarithmic or semilogarithmic axes. These fits are plotted along with the data used to produce them in Fig. 10. The slight difference from \(-1 \) in the scaling exponent for \( \gamma^{(h)}_c \) is due to numerical finite-size effects.

<table>
<thead>
<tr>
<th>Quantity</th>
<th>Scaling</th>
<th>( 1 - r^2 )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( P^{(QW)}_{\text{max}} )</td>
<td>( 1 - 1.734n^{-1.112} )</td>
<td>( 7.820 \times 10^{-5} )</td>
</tr>
<tr>
<td>(</td>
<td>E_0</td>
<td>^2 +</td>
</tr>
<tr>
<td>(</td>
<td>\psi_{\text{null}}</td>
<td>E_0</td>
</tr>
<tr>
<td>( \gamma^{(h)}_c )</td>
<td>( 1 - 1.233n^{-0.0425} )</td>
<td>( 1.120 \times 10^{-5} )</td>
</tr>
</tbody>
</table>

IV. HYBRID ANNEALING SCHEDULES

Having arrived at a common Hamiltonian form for QW and AQC searching on an \( n \)-dimensional hypercube and having derived optimal coefficients for each case, we can now interpolate the coefficients to generate hybrid search algorithms. In this section we show how this may be done and study the resulting dynamics. We begin by looking at small systems with \( n = 5 \) and \( 8 \) and then study the dynamics of systems with very large \( n \) by demonstrating that this limit corresponds to a two-state single-avoided-crossing model.

A. Motivation and definition

We have already noted that QW and AQC search algorithms both use the same terms in the Hamiltonian, differing only in the time dependence. With appropriate choice of parameters, both provide a quadratic quantum speedup: a search time proportional to \( \sqrt{N} \) for a search space of size \( N \). This suggests the question of whether we can map smoothly between QW and AQC searching while maintaining the quantum speedup.

To construct the mapping, we generalize the AQC Hamiltonian of Eq. (11) by defining a time-dependent Hamiltonian

\[
\hat{H}(\tau) = A(\tau)\hat{H}_0 + B(\tau)\hat{H}_p
\]

as a function of the reduced time \( \tau = t/\tau_f \), where the annealing schedules \( A(\tau) \) and \( B(\tau) \) satisfy \( A(0) \gg B(0) \) and \( B(1) \gg A(1) \). The AQC algorithm as described by Eq. (11) is obtained by setting

\[
A_{\text{AQC}}(\tau) = 1 - s(\tau),
\]

\[
B_{\text{AQC}}(\tau) = s(\tau).
\]

The QW search Hamiltonian with hopping rate \( \gamma \), described by Eq. (9), can also be obtained by setting

\[
A^{(QW)}_{\gamma}(\tau) = \begin{cases} 
\gamma, & \tau < 1 \\
0, & \tau = 1,
\end{cases}
\]

\[
B^{(QW)}_{\gamma}(\tau) = \begin{cases} 
1, & \tau > 0 \\
0, & \tau = 0.
\end{cases}
\]

We can make this even closer to the AQC form by defining \( \beta = 1/(1 + \gamma) \) and setting

\[
A_{\text{QW}}(\tau) = \begin{cases} 
1 - \beta, & \tau < 1 \\
0, & \tau = 1,
\end{cases}
\]

\[
B_{\text{QW}}(\tau) = \begin{cases} 
\beta, & \tau > 0 \\
0, & \tau = 0.
\end{cases}
\]

For a QW search on the hypercube, using Eq. (10) for \( \gamma^{(h)}_c \), to achieve optimal \( \sqrt{N} \) scaling we must set \( \beta \) equal to

\[
\beta^{(h)}_c = \frac{1}{1 + R_1}.
\]

For \( 0 < \tau < 1 \), the reparametrization of Eq. (27) in Eq. (28) maintains the ratio of \( A_{\text{QW}}(\tau)/B_{\text{QW}}(\tau) = \gamma \). However, it also introduces a global energy shift \( A_{\text{QW}}(\tau) = \beta A^{(QW)}_{\gamma}(\tau) \) and \( B_{\text{QW}}(\tau) = \beta B^{(QW)}_{\gamma}(\tau) \). The observant reader will note that, because the optimal \( \gamma^{(h)}_c \) is dependent on the size of the system, this reparametrization introduces a weak dependence of the global energy scale on system size \( N = 2^n \). However, since \( \beta^{(h)}_c \rightarrow 1 \) in the large-\( N \) limit, this weak dependence cannot affect the leading-order term in the asymptotic scaling and the reparametrized quantum-walk-search algorithm still provides optimal \( \sqrt{N} \) scaling.
In the way we have parametrized them above, the AQC and QW protocols differ only in the annealing schedules $A(\tau)$ and $B(\tau)$. Hence, we can use the QW and AQC schedules as end points of a smooth interpolation between these two search algorithms to define a continuum of hybrid protocols. Using a parameter $\alpha \in [0, 1]$, where $\alpha = 0$ corresponds to a quantum walk and $\alpha = 1$ corresponds to AQC, we can define

$$A(\alpha, \beta, \tau) = \frac{1 - s(\tau)}{\alpha + (1 - \alpha)\frac{(1 - \beta)}{(1 - \beta)}}$$

$$B(\alpha, \beta, \tau) = \frac{s(\tau)}{\alpha + (1 - \alpha)\frac{(1 - \beta)}{(1 - \beta)}} \tag{30}$$

giving a family of hybrid quantum algorithms defined by the Hamiltonian

$$\hat{H}_{AB} = A(\alpha, \beta, \tau)\hat{H}_0 + B(\alpha, \beta, \tau)\hat{H}_p. \tag{31}$$

This interpolation is quite general, for well-behaved $\hat{H}_0$ and $\hat{H}_p$, with the caveat about the extra dependence of the energy scale on the QW hopping rate through $\beta$ mentioned above. The resulting family of functions is illustrated in Fig. 4 for search over five- and eight-qubit hypercube graphs.

Note that, although it does not follow a priori from the construction that these interpolated AQC-QW schedules will yield a quantum speedup at all for searching, let alone an optimal $\sqrt{N}$ scaling. This is because the different mechanisms in a quantum walk and AQC could be incompatible in combination. We return to this important question in Sec. IV C, where we show that properly specified interpolations can indeed achieve the theoretical optimum $\sqrt{N}$ scaling.

### B. Small-size examples

To gain intuition for how our interpolated schedules behave, we study small systems of five and eight qubits. These have been simulated using the full Hamiltonian on the hypercube; for numerical methods see Appendix B. Figure 5 shows how the final success probability varies with the search duration $t_f$ for a QW, an AQC, and an intermediate $\alpha = 0.5$ search over the five-qubit hypercube graph. Note that, because the schedules $A$ and $B$ are in general nonlinear functions of time, in all plots against $t_f$, each point represents a separate run of the quantum search algorithm for that value of $t_f$; the plots do not also represent the time evolution $0 \leq t \leq t_f$, except for $\alpha = 0$ when the schedule functions are constant ($A = 1 - \beta$ and $B = \beta$). Plots of the time evolution for a single search can be seen in Sec. V C. Also plotted in Fig. 5 are the annealing schedules $A$ and $B$ as a function of the reduced time $\tau$, illustrating how the shape of the functions $A(\alpha, \tau)$ and $B(\alpha, \tau)$ changes for different values of $\alpha$, from flat for a quantum walk to a curving AQC annealing schedule for $\alpha = 1$. We see that the qualitative behavior of adiabatic evolution is fundamentally different from that of the quantum-walk search. For the optimal AQC schedule the success probability increases monotonically to a value very close to one. In contrast, a quantum walk shows oscillatory behavior, and although the success probability does not approach one, it does show a faster initial increase than for AQC. The intermediate schedule shows a mix of both behaviors, with a locally oscillating but globally increasing success probability that shows an initial increase rate between that of a quantum walk and AQC. For a continuum of $\alpha$ values, Figs. 6 and 7 illustrate the same qualitative behavior for five-qubit and eight-qubit quantum searches. The oscillatory behavior associated with a quantum walk slowly fades away as the interpolation approaches the respective AQC schedule, at which point the success probability $P$ increases monotonically with $t_f$. If a relatively low success probability is sufficient, only a short total runtime $t_f$ is needed and a quantum walk is the best strategy. As $t_f$ is increased, the best strategy is to increase $\alpha$ and start...
given runtime $t_f$ and value of $\alpha$. Figure 7 shows the same comparisons for the slightly larger value of $n = 8$ qubits. The optimal $\alpha$ moves away from $\alpha = 0$ at a smaller value of $t_f$ and $P$ for $s^{(c)}$ than it does for $s^{(n)}$. There is also more structure in the optimal $\alpha$ line (black dashes) for $s^{(c)}$ than for $s^{(n)}$, with a range of $\alpha$ values that are optimal for more than one value of $P$. Otherwise, the two behave quite similarly for these small sizes, suggesting that both $s^{(c)}$ and $s^{(n)}$ are able to provide a quantum speedup for hybrid protocols. To confirm this in general, not just for small $n$, further analysis and simulations of larger systems are required, which we tackle in the following sections.

### C. Performance of hybrid algorithms

Our strategy for analyzing the scaling of the hybrid quantum search algorithms is to show that the performance is dominated by a single, low-energy, avoided crossing (see Fig. 1), which is present at the same position in all our hybrid algorithms. We then show that the essential features of the behavior are captured by a simple two-state single-avoided-crossing model which all the hybrid algorithms map to in the large-size limit. For this simple avoided-crossing model we can easily show that the hybrid algorithms all provide an optimal quantum speedup. It then follows that our full-size hybrid algorithms have the same asymptotic scaling.

We first consider the end points of the interpolation, QW and AQC searches. For the AQC search, the optimal schedule $s^{(c)}(\tau)$ or $s^{(n)}(\tau)$ is derived directly from the functional form of the lowest avoided crossing, ensuring that the Hamiltonian is changed slowly enough to avoid transitions to higher-energy levels. We only need to show that the low-energy structure of the Hamiltonian is dominated by a single avoided crossing throughout the process. This is shown numerically in Fig. 8. The width $w(n)$ of the avoided crossing decreases rapidly with $n$. Even for a modest size of $n = 50$ qubits, the switch from 95% overlap with the hypercube Hamiltonian ground state to 95% overlap with the marked state occurs in less than $10^{-8}$ of the total dynamic range of the protocol, which runs from $s(\tau) = 0$ to $s(\tau) = 1$. In contrast, for the QW search, transitions to higher-energy levels are a necessary part of the evolution to the marked state, so we need to determine the scaling of several related quantities to show that a single avoided crossing dominates in determining the behavior.

### D. Minimum gap scaling in the QW search

For the QW search, to show numerically that the lowest avoided crossing is the only relevant feature in the large-$N$ limit, we must demonstrate two things: first, that the minimum gap $g_{\text{min}} = (E_1 - E_0)$ between the ground state and the first

---

1While the dynamical range $w(n) = \Delta s(\tau)$ in which the state rotates between the two nearly orthogonal states $|\psi_{\text{init}}\rangle$ and $|m\rangle$ becomes exponentially small, the total runtime $t_f \sim \sqrt{N} = \exp(n/2)$ grows even more quickly, so the time taken $w(n)/t_f$ becomes exponentially small as $n \rightarrow N$. But for $w(n)/t_f$ there is no exponential dependence on $n$.
excited state becomes much smaller than the minimum gap between the ground state and the second excited state, and second, that the lowest avoided level crossing, where \( g(\tau) = g_{\text{min}} \), dominates the transition between the ground state of the hypercube Hamiltonian \( \hat{H}_0 \) and the ground state of the marked state Hamiltonian \( \hat{H}_e \) becomes more dominant as system size increases. Noting that, as illustrated in Fig. 4, around the minimum gap, where all the schedules cross, we have \( 1 - s(\tau) \approx \gamma^{(h)}_{\text{min}} \), Fig. 8 shows that both of these do in fact occur. The left inset shows that at the avoided crossing, the gap between the ground state and first excited state shrinks exponentially faster in \( n \) than the gap between the ground state and second excited state. The main figure and right inset of Fig. 8 show how the transition between the two ground states becomes dominated by the dynamics at \( g_{\text{min}} \) as \( n \) increases.

For a pure quantum-walk search, this convergence to behavior dominated by a single avoided crossing can be seen in Fig. 9, which shows not only that does the search success probability \( P \) approach one in the large system limit, but also that the time evolution of \( P \) (shown in the inset) approaches the functional form for the single avoided crossing \( P(\tau) = \sin^2(\gamma_{\text{min}} \tau/2) \). The nonsinusoidal shapes of these curves at low qubit number are due to the influence of excited states higher than the first exited state. These small-size effects are clearly significant up to about \( n = 12 \) qubits. This highlights the potentially atypical nature of the five- and eight-qubit examples in Sec. IV B and the importance of examining larger system sizes. For \( n > 12 \), the probability \( P \) smoothly approaches one, although relatively slowly (polynomially) as a function of \( n \). Based on the data in Table I, we can deduce that this effect relates to the fact that the overlap of the manifold
The initial state |1⟩ is only an approximate eigenstate of $\hat{H}_0^{(AC)}$, but the approximation improves as $g_{min}$ decreases. Solving the eigensystem for this Hamiltonian gives

$$g^{(AC)}(s) = \left\{ (1 - 2s)^2 + 4g_{min}(1 - s^2) \right\}^{1/2}$$

for the gap between the two energy levels. In the limit of small $g_{min}$ the minimum gap is $g_{min}$ and occurs for $s = \frac{1}{2}$. We can then apply the method of [21] to find the optimal schedule $s(t)$ for this system. Calculating $dH/ds$, we find

$$\frac{dH^{(AC)}}{ds} = -\hat{\sigma}_z + g_{min}\hat{\sigma}_x,$$

(35)

giving a maximum value of one\(^2\) for $\langle (dH^{(AC)}/ds)_{0,1} \rangle$ in the large-size limit. Using Eq. (20) to find the optimal schedule, we need to solve

$$\frac{ds}{dt} = \epsilon g^{(AC)}(s) \frac{1}{g_{min}} = \epsilon \left\{ (1 - 2s)^2 + 4g_{min}(1 - s^2) \right\},$$

(36)

where the maximum value is used for $\langle (dH^{(AC)}/ds)_{0,1} \rangle$. This can be integrated straightforwardly to give

$$\arctan \left( \frac{2g_{min}(s - 1) + 2s - 1}{g_{min}} \right) = 2g_{min}\epsilon t + c,$$

(37)

with

$$c = -\arctan \left( \frac{2g_{min} + 1}{g_{min}} \right).$$

(38)

From this we find for $s = 1$ that the runtime $t_f^{(AC)}$ is given by

$$\epsilon t_f^{(AC)} = \frac{\pi}{2g_{min}} \arctan(1/g_{min}) \approx \frac{\pi}{2g_{min}} - 1,$$

(39)

where the approximate expression uses $\arctan(1/g_{min}) \approx \frac{\pi}{2} - g_{min}$ for $g_{min} \ll 1$ and terms of order $g_{min}^2$ have been dropped. The runtime of the optimal schedule thus depends inversely on the size of the minimum gap, as expected. Solving for $s(t)$ and dropping terms of order $g_{min}^2$ gives

$$s(t) \approx \frac{1}{2} [1 - g_{min}\cot(g_{min}(2\epsilon t + 1))].$$

(40)

In this limit where $g_{min} \ll 1$, an equivalent way to parametrize $H^{(AC)}$ is

$$\hat{H}^{(AC)} = \frac{g_{min}}{2} [f(t)\hat{\sigma}_z - \hat{\sigma}_x],$$

(41)

where $-\infty < f(t) < \infty$. This form is obtained by taking $(1 - 2s(t))/g_{min} \rightarrow f(t)$ and shifting the zero point of the energy scale to the middle of the avoided crossing. As $f(t)$ changes from $-\infty$ to $\infty$ it passes through zero as the sign of the $\hat{\sigma}_z$ term changes, when the $\hat{\sigma}_x$ term drives the transition from $|1⟩$ to $|0⟩$. Although the $\hat{\sigma}_z$ term is no longer turned off at the end of the schedule, it becomes negligible in comparison to the $\hat{\sigma}_x$ term and does not significantly alter the dynamics. This can be intuitively thought of as scaling all features of $H^{(AC)}$ other than the avoided crossing to $\pm \infty$.

\(^2\)Strictly the maximum value is $\sqrt{1 + g_{min}^2} = 1 + O(g_{min}^2)$, however this correction simply modifies $\epsilon$ in what follows and disappears altogether when terms of order $g_{min}^2$ are dropped.
The QW form of the single-avoided-crossing search Hamiltonian is also simple to analyze. We deduce the optimal value of $\gamma_0 = 1$ from the value of $s = \frac{1}{2}$ at the avoided crossing. We then use Eqs. (28) in which $\beta_0 = 1/(1 + \gamma_0) = \frac{1}{2}$, whence

$$H_{\text{QW}}^{(AC)} = (1 - \beta_0)H_{0}^{(AC)} + \beta_0 H_{p}^{(AC)}$$

$$= \frac{1}{2} \left\{ \left( 1 + \delta_t \right) - g_{\text{min}} \delta_t + \frac{1}{2} (\mathbb{1} - \delta_t) \right\}$$

$$= \frac{1}{2} (1 - g_{\text{min}} \delta_t). \quad (42)$$

The $\delta_t$ term causes deterministic transitions between the two states regardless of their energies, at a rate determined by $g_{\text{min}}$. By solving for the dynamics, the time for the input state $|1\rangle$ to evolve to the marked state $|0\rangle$ can be shown to be $t_{f}^{(\text{QW})} = \pi / g_{\text{min}}$.

We can now map between QW and AQC in the avoided-crossing model using Eqs. (30) for $A(\alpha, \beta, \tau)$ and $B(\alpha, \beta, \tau)$. Using $\beta = \frac{1}{2} = 1/(1 + \gamma_0)$, for $s(t)$ from Eq. (40) we have hybrid schedules

$$A_{\text{AC}}(\alpha, t) = \frac{1 - s(t)}{\alpha + 2(1 - \alpha)[1 - s(t)]}.$$  

$$B_{\text{AC}}(\alpha, t) = \frac{s(t)}{\alpha + 2(1 - \alpha)s(t)} \quad (43)$$

We can easily show numerically that all the hybrid algorithms defined by Eqs. (43) find the marked state with high probability (given by $\epsilon$) in a runtime $\lesssim t_{f}^{(\text{AC})}$ given by Eq. (39), the runtime required by the optimal AQC $s(t)$ used to define the hybrid schedules. Figure 11 shows this is indeed the case. The white contours highlight the difference between the pure QW search, which succeeds with certainty, and the AQC and hybrid algorithms, which always have a probability of error $\epsilon^2$ that can be traded against the runtime $t_f$. The shallow upward curve of these contours towards the AQC end of the hybrid protocols shows in what sense the QW search is better than AQC in the large-size limit.

The hybrid algorithms on the full hypercube map onto the hybrid single-avoided-crossing-model algorithms for large $n$.

This follows from the solution methods for the end points, QW and AQC searches, which all use the two-level approximation to prove the quadratic speedup. Since the full hypercube hybrid algorithms are defined from these in the same way as the single-avoided-crossing-model hybrid algorithms are defined, the hybrid algorithms also map to the corresponding single-avoided-crossing hybrid algorithm. They therefore also obtain the quantum speedup for large $n$, which is what we set out to show.

F. Optimal hybrid algorithm for a single run

Having shown that hybrid protocols between QW and AQC searching maintain the quadratic quantum speedup, the next question is how to optimize over this continuum of hybrid schedules for finite-size systems. The single-avoided-crossing model gives the large-size limit in which QW protocol is the optimal strategy. However, this limit is only reached in a polynomial scaling with $n$, as described in Sec. IV D.

For a single run of a search algorithm, we can trade off between the magnitude of the success probability and the runtime of the search. For QW searches, there is a maximum probability $P_{\text{max}}^{(\text{QW})}$ that can be obtained; shorter runtimes reach lower success probabilities and so do longer runtimes. For AQC searches a longer runtime always reaches a higher success probability. We can thus specify the success probability we require and ask which hybrid algorithm attains this success probability with the shortest runtime. We consider multiple run strategies in Sec. V.

As Fig. 12(a) illustrates for sizes from $n = 12$ to $n = 20$, the optimal protocol jumps from QW to hybrid at $P \approx P_{\text{max}}^{(\text{QW})}$ and the optimal hybrid strategy it jumps to becomes more QW-like (smaller $\alpha$) as the system size increases. As $P$ is
increased further, the optimal hybrid strategy becomes steadily more AQC-like (larger $\alpha$). Figure 12(b) shows that the hybrid strategies require runtimes $t_f$ larger than $g_{\text{min}}/\pi$ to achieve higher success probabilities in a single run.

V. MULTIPLE RUNS FOR ONE SEARCH

In the previous sections we derived hybrid search Hamiltonians for the hypercube and studied their dynamics. However, this does not yet give us a full picture of the relative usefulness of the different dynamics. In this section we study the relative performance of the different searches when we allow for the possibility of multiple searches and when the system suffers from decohering interactions with its environment.

A. Motivation

In a realistic setting of the search problem we can easily check whether the result of a search is the correct answer or not. Hence, we must consider not only single-run strategies, but also multirun strategies, where the success probability is defined as the probability of succeeding in at least one of several runs. In the context of a quantum search on the hypercube, we measure which site of the hypercube our state is on and then determine the energy of this state with respect to the search Hamiltonian. If this energy is zero, then we have found the state we are looking for; otherwise we would need to account for a nonzero initialization time $t_{\text{init}}$ associated with each run of the search. Such an initialization time is mathematically as well as physically necessary. The fidelity between the initial state and marked state $|\langle \psi_{\text{init}} | m \rangle|^2 = \frac{1}{N}$ is nonzero. An arbitrarily short run is equivalent to making a random guess. Therefore, without an additional penalty per run, it would be possible to guess an arbitrarily large number of times for free, thus finding the marked state in a total arbitrarily short time. Any physical device will take a significant amount of time both to setup the initial state and to measure the final state. For the purposes of our study, the effects on the total search time of initialization and readout times are the same, therefore the quantity we call $t_{\text{init}}$ should be taken to include all of the time associated with a single run other than the actual runtime of the algorithm $t_f$, i.e., as including both initialization and measurement.

B. Multiple-run searching

As examples, we consider $n=12$ and 14 qubits using the numerically calculated optimal strategy $s^{(n)}$. Referring to Fig. 9, $n=12$ still shows finite-size effects, while $n=14$ is just into the smoothly scaling regime. We find for chosen success probabilities in the range 0.95–0.99 that the optimal strategy depends on both $t_{\text{init}}$ and $P_{\text{target}}$ as shown in Fig. 13. For the range of $t_{\text{init}}$ we examine, both sizes show a transition from a single run able to reach the required success probability to a region requiring two runs. The single runs are hybrid, becoming progressively more AQC-like as the target success probability increases further. Finite-size effects are visible for $n=12$ in the nonmonotonic shape of the boundary between one run and two runs in Figs. 13(a) and 13(c). For smaller $n < 12$, these effects become more complicated; there is no single best strategy for a small search space. Indeed, we also found that the optimal strategy changes significantly when any of the parameters are varied. The complexity in the optimal search strategy for small $n$ is because the two-level approximation does not hold well in this regime and interactions with higher excited states have a non-negligible effect. This suggests that a similarly complex situation will likely be present in more sophisticated optimization Hamiltonians, whenever a two-level approximation is not valid.

C. Noisy quantum searching

Another realistic situation where multiple runs can be helpful is when there is a significant level of unwanted decoherence or other forms of noise acting on the quantum hardware. In this case, shorter runs that end before decoherence effects are too strong, but consequently have lower success probabilities and hence need more repeats, may be able to maintain a quantum speedup. Decoherence effects on the different AQC and QW mechanisms are analyzed in more detail in related work and the effects of noise in AQC search have been studied in [44,45]. Here we focus on hybrid algorithms and the extra options these provide for optimizing the search.

We choose a simple model of decoherence by adding a Lindblad term to the von Neumann equation for the system density operator $\hat{\rho}(t)$,

$$\frac{\partial \hat{\rho}(t)}{\partial t} = -\frac{i}{\hbar} [\hat{H}(t), \hat{\rho}(t)] + \kappa P[\hat{\rho}(t)],$$

where $\hat{H}(t)$ is the search Hamiltonian and $\kappa P[\hat{\rho}(t)]$ is a decoherence term tuned by a rate $\kappa$. We choose a form for $P$ that uniformly reduces the coherences between states corresponding to vertices of the hypercube (the computational basis). This type of decoherence has been well studied in the context of quantum walks [46–48] and, for high decoherence rate $\kappa \gg \gamma$, can be thought of as continuous measurement in

\[022339-13]
the search space resulting in a quantum Zeno effect [49]. It is equivalent to coupling with an infinite-temperature bath.

Since we now have five parameters to optimize over for a given search size \( n \) (\( P, t_f, \alpha, \kappa, \) and number of runs \( r \)), we first consider single-run searches with success probability \( P(t_f, \alpha, \kappa) \). This is the final success probability of a hybrid search specified by \( \alpha \) of duration \( t_f \) in the decoherence model of Eq. (44) with decoherence rate \( \kappa \). We simulate the searches for durations \( 0 \leq t_f \leq 200 \) and define the search duration \( t_o \) that maximizes \( P \) for a particular choice of \( \alpha \) and \( \kappa \). We also define \( \alpha_o \) as the value of \( \alpha \) which maximizes \( P(t_o, \kappa, \alpha) \), which corresponds to the search that reaches the highest success probability for a given decoherence rate \( \kappa \). Note that, for computational reasons, we limit \( \alpha \) to the values 0.0, 0.1, 0.2, \ldots, 0.9, 1.0 when performing the maximizations; intermediate values are of course possible.

We begin by looking at how the instantaneous success probability \( P(t) = \langle m | \rho(t) | m \rangle \) evolves during a search, where \( m \) denotes the marked site. Inset in Fig. 14 are plots of the evolution of \( P \) during a search over a seven-qubit hypercube graph for varying decoherence rates \( \kappa \), in terms of reduced time \( \tau = t/t_f \), for a \( t_f \) that shows the first peak of the QW search. The broad effect of the decoherence is to reduce the instantaneous success probability towards a value of \( 1/N \), equivalent to classical guessing. The QW, AQC, and hybrid search algorithms retain their characteristics up to an overall decoherence damping, which is independent of \( \alpha \). As can be seen for the \( \kappa = 0 \) subplot (left) in Fig. 14, the QW search spreads out more quickly over the search space and therefore exhibits a more rapid initial increase in \( P \). On the other hand, AQC searching can reach higher values of \( P \) for sufficiently small values of \( \kappa \), albeit at later times.

Figure 14 shows which is fastest out of individual QW, AQC, and hybrid searches for a single search, for a given value of \( \kappa \) and of \( P \); from AQC through to the hybrid when maximal success probability is required, with the QW search performing best for slightly lower values of \( P \). This indicates a remarkably large range of situations where QW dynamics is desirable, either as part of hybrid algorithms that hit the highest success probabilities for all but the smallest decoherence rates or alone in the form of a static Hamiltonian, if a marginally smaller success probability can be tolerated.

Another way to compare the different searches under decoherence is to ask whether a QW, AQC, or hybrid search will give the maximum possible success probability \( P \) for a given value \( \kappa \). Figure 15(b) shows how this maximum \( P \) varies for the three cases, as well as the value of interpolation parameter \( \alpha_o \) for the best-case hybrid search. For small values of \( \kappa \), \( \alpha_o = 1 \), i.e., AQC gives the highest peak success probability. As \( \kappa \) is increased, the highest-scoring search changes and \( \alpha_o \) decreases monotonically, indicating hybrid searches perform the best overall for intermediate levels of decoherence. In the limit of very high decoherence we are in a quantum Zeno effect regime which keeps the search in the initial superposition over all possible states. This means all searches will succeed with the same probability \( P = |\langle \psi_{\text{final}} | m \rangle |^2 = 1/N \), equivalent to classical guessing. The usefulness of a search is also determined by how quickly it can be performed, and so the search time \( t_f \) is shown in Fig. 15(a), showing that while the QW search never has the highest success probability in the range we examine, it can be substantially quicker. This helps to explain why hybrid schedules take on more QW character as \( \kappa \) is increased and soon begin to achieve higher success probabilities than AQC in shorter search times.

Having characterized the effects of decoherence on a single run, we now consider multiple-run search strategies where each search is of the same duration \( t_f \). We define the optimal

\[ P \]
the region of instantaneous measuring with $\tau_f$ and/or target success probability $P$ have increased such that the best performing strategy with $r$ searches drops below $P$, and another run is required. In this case the target can be reached by $r + 1$ lower quality searches. This drop in the quality required of the single search means that a faster more QW-like search can be used to succeed and therefore the optimal value of $\alpha$ drops.

Our numerical results for hybrid algorithms in the presence of noise can be understood intuitively by considering how $P$ scales with a small amount of noise in the AQC and QW edge cases. For noise rate $\kappa$ per unit time, the success probability for a single run reduces as $P \approx \exp(-\kappa t_f)$, where $t_f$ is the time taken for one run of the search algorithm. For $P \sim 1$ we thus require $\kappa t_f \ll 1$, i.e., $\kappa \ll 1/t_f$. For QW searching on the hypercube, we have $t_f^{(\text{QW})} \approx \frac{2}{\sqrt{N}}$, hence we obtain $\kappa_{\text{QW}} \ll 2/\pi \sqrt{N}$ for tolerable noise rates. For AQC, on the other hand, from Eq. (24) we have $t_f^{(\text{AQC})} \approx \frac{4}{\sqrt{N}}$ for large $N$. For high success probability, since $P \sim 1 - \epsilon$, the adiabatic condition requires $\epsilon \ll 1$ and we obtain $\kappa_{\text{AQC}} \ll 4\epsilon / \pi \sqrt{N}$. The extra factor of $\epsilon$ implies $\kappa_{\text{AQC}} \ll \kappa_{\text{QW}}$. Hence, the QW search will be more robust to disturbance by noise, as we have found numerically for the single-run case. For our $n = 7$ example, $\kappa_{\text{QW}} \ll 0.056$ and $\kappa_{\text{AQC}} \ll 0.11 \leq 0.011$ for $P = 0.99$ and indeed we see in Fig. 14 that performance drops below $P = 0.5$ for $\kappa_{\text{QW}} \geq 0.025$. However, when multiple runs are included, hybrid strategies with significant adiabatic character can still outperform the QW search, depending on hardware characteristics determining the initialization and measurement time required per run.

VI. PROBLEM MISSPECIFICATION

So far we have studied the dynamics of the hybrid search Hamiltonians as part of single- and multiple-run algorithms and in the presence of noise. In the following section we consider misspecification of the problem, for which the dynamics remain coherent, but some parameters are changed in unknown ways.

A. Motivation

Studying the effects of problem misspecification is particularly relevant given the critical difficulties which many classical analog computing efforts have faced due to propagation of errors [50]. Misspecifications can come about in a variety of ways, such as limited precision for setting the controls in the computer, ignorance of what the optimal parameters should be, or noise which is at a much lower frequency than the rate of the relevant quantum dynamics. An important example of the latter is so-called $1/f$ noise in superconducting qubit devices [51,52], such as the quantum annealers constructed by D-Wave Systems Inc. It has been shown, for instance, that such misspecifications can cause AQC to give an incorrect solution on Ising spin systems [53,54] and it effectively limits the maximum useful size of such devices. For an example of

FIG. 16. Optimal hybrid search parameter $\alpha$, and number of runs $r$, for multiple-run searching on an $n = 7$ hypercube. The optimal search is that which achieves the target success probability $P$ in the shortest total time $t_f(t_f + t_{\text{init}})$, where $t_{\text{init}}$ and $t_f$ are the initialization time and runtime, respectively. (a) and (b) Dependence on $t_{\text{init}}$ and decoherence rate $\kappa$ when $P$ is set equal to 0.95. The black indicates the region of instantaneous measuring with $t_f = 0$, where $r = 582$. (c) and (d) Dependence on $P$ and $\kappa$ when $t_{\text{init}}$ is set equal to 10. Plots (a) and (c) show $\alpha$, and (b) and (d) show $r$. The analytic expression (21) is used for the AQC schedule. Time and rate units are given by Eq. (25).

The boundaries where another run is required correspond exactly to the regions where the optimal value of $\alpha$ suddenly drops. This transition arises when the decoherence rate $\kappa$ and/or target success probability $P$ have increased such that the best performing strategy with $r$ searches drops below $P$, and another run is required. In this case the target can be reached by $r + 1$ lower quality searches. This drop in the quality required of the single search means that a faster more QW-like search can be used to succeed and therefore the optimal value of $\alpha$ drops.

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There is a small threshold initialization time below which the best strategy is to take multiple measurements of the system state as soon as it is prepared at a small cost $t_{\text{init}}$, indicating that our device can do no better than classical random guessing. Other than this threshold, there is little dependence on initialization time. There is a broad tendency towards AQC-like searches as $P$ is increased, however for larger values of $\kappa$ an AQC search ceases to ever be optimal and hybrid or QW searches are preferred. As $\kappa$ is increased, there is a localized trend for more AQC-like searches to be optimal, however this is punctuated with discontinuous changes to a more QW-like search. The reason for these discontinuous changes can be seen in Figs. 16(b) and 16(d).
the effects of problem misspecification on a real experiment, see [14].

For this work, we will consider simple misspecification models in the large-system limit, where the Hamiltonian can be mapped to a single avoided crossing in the form of Eq. (33) or (41). For the purpose of studying problem misspecification, it is most convenient to work with the form in Eq. (41), which we use for the duration of this section. Because the initial and marked states are orthogonal in this limit, considering multiple runs which can be performed with negligible initialization and readout time, the new success probability for a given anneal time can be modeled as an uncertainty in the total energy scale through Eq. (41). The effect of uncertainty can be seen in Fig. 11. (a) Probability $P$ of finding the marked state versus runtime and $\alpha$ for the single-avoided-crossing model, same as in Fig. 11. (b) Same as in (a) with a 30% misspecification of the energy $\Delta g_{\min}$.

B. Error in gap size

The effect of misspecifying the size of the minimum gap can be modeled as an uncertainty in the total energy scale $\Delta g_{\min}$, which is equivalent to a misspecification of the total runtime $t_f$ through Eq. (41). The effect of uncertainty can be modeled by performing a convolution of the success probability versus runtime with a distribution describing the uncertainty. An example result of such a convolution is depicted in Fig. 17(b). Assuming that the misspecification is distributed in a Gaussian manner around the intended runtime, the new success probability for a given anneal time $t_f$ and $\alpha$ becomes

$$P(t_f, \alpha, \Delta g_{\min}) = \int_{-\infty}^{\infty} dt' \frac{P(t'_f, \alpha)}{\Delta g_{\min}} \exp \left\{ -\frac{(t'_f - t_f)^2}{2(\Delta g_{\min})^2} \right\},$$

where $\Delta g_{\min}$ is the (unitless) fractional uncertainty in $g_{\min}$ and the absolute value in the argument of $P$ within the integral is included to avoid negative time arguments. For reasonable values of $\Delta g_{\min}$, it will be rare for $t'_f < 0$ and the effect of taking the absolute value will be negligible.

Figure 17 shows how the evolution makes a smooth transition between the characteristically sinusoidal behavior of success probability versus runtime for a quantum walk and the characteristically monotonic behavior of AQC. As the comparison between the perfect and misspecified cases demonstrates, gap misspecification causes a large reduction in the success probability of QW protocols, but has almost no effect on the monotonic AQC search. Figure 18 illustrates that, for moderately high success probability and moderate amounts of misspecification of $\Delta g_{\min}$, the best protocol is no longer a quantum walk, but lies in between the optimal AQC schedule and a quantum walk. For large gap misspecification where a high success probability is required, the best approach is to run an intermediate strategy twice.

The reason that gap-size misspecification makes hybrid protocols ($\alpha > 0$) outperform QW protocols for a large range of parameter space is because a quantum walk can only succeed with a probability approaching one if $t_f R_{\min}$ is an odd multiple of $\pi$. The misspecification smears out these peaks and implies that the success probability of a quantum walk will not approach one for any value of $t_f$. For protocols with some adiabatic character, however, the maximum success probability will still approach one as $t_f$ becomes larger, as the adiabatic theorem holds for any finite gap. In cases where the misspecification overstates the size of the gap the success probability of AQC will actually improve.

C. Error in avoided-crossing location

Another type of problem misspecification is incorrectly specifying the position of the avoided crossing. To model this, we consider a modification of the problem Hamiltonian

$$\hat{H}^{(AC)}(t, q) = \hat{H}^{(AC)}(t) + q \frac{g_{\min}}{2} \hat{z},$$

where $\Delta g_{\min}$ is the (unitless) fractional uncertainty in $g_{\min}$ and the absolute value in the argument of $P$ within the integral is included to avoid negative time arguments. For reasonable values of $\Delta g_{\min}$, it will be rare for $t'_f < 0$ and the effect of taking the absolute value will be negligible.

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This addition to the problem Hamiltonian provides a shift in the avoided-crossing position \( f(t) \rightarrow f(t) + q \) in Eq. (41). Effectively introducing this shift causes the schedule to slow down at the wrong point, reducing the success probability. As we did for the case of gap mis-specification, we can model the effect of this error as a convolution of the success probability distribution with \( q \) with a Gaussian of width \( \Delta q \). We define the success probability with misspecified avoided-crossing position as

\[
P(t, \alpha, \Delta q) = \int_{-\infty}^{\infty} dq \frac{P(t, \alpha, q)}{\Delta q \sqrt{2\pi}} \exp \left( -\frac{q^2}{2(\Delta q)^2} \right). \tag{47}
\]

where \( \Delta q \) is the (unitless) fractional uncertainty in \( q \) that controls the degree of misspecification. Figure 19 illustrates that, in contrast to gap mis-specification, the best strategy is almost always a quantum walk. Intermediate strategies only become the superior method briefly, at the edge of the regime where single runs are the best way to reach the desired probability. At higher misspecification, multiple repeated quantum walks become the best strategy.

Misspecification in the avoided-crossing position does significant harm to both AQC and QW protocols. The success probability of a QW protocol performed with an incorrectly chosen \( \gamma \) does not approach one. Similarly, an AQC protocol with a poorly chosen schedule will require a much longer runtime for the success probability to approach one. The faster runtime of the quantum walk then means it beats AQC for multiple runs.

VII. SUMMARY AND OUTLOOK

In this paper we provided a detailed study of the scaling of continuous-time quantum search algorithms on a hypercube graph. Noting that both quantum-walk and adiabatic quantum search algorithms can be expressed as two extremes of quantum annealing schedules, we defined a family of quantum search algorithms that are hybrids between QW and AQC algorithms. By mapping the algorithms to a one-qubit single-avoided-crossing model, we showed that the whole family achieves the maximum possible \( \sqrt{N} \) quantum speedup. There are a number of subtleties in the scaling behavior on the hypercube that we treated in detail for short search times, complementing the work by Weibe and Babcock [42] on long timescales.

Our hybrid QW-AQC schedules are an example of the advantages we gain by treating both a quantum walk and AQC as part of the same method of continuous-time quantum computing. We found that hybrid strategies intermediate between a quantum walk and AQC provide the best quantum search algorithm under a range of realistic conditions. The techniques we used here can easily be extended to a hybrid quantum search on other graphs and to other quantum-walk or adiabatic quantum computing algorithms.

This work focused on the search problem due to its relative simplicity and the fact that annealing schedules can be derived analytically, which we do for the hypercube graph in Appendix A. The core ideas and methods are quite general and can easily be extended to more complex and realistic problems, such as a fixed-point search, where multiple states are marked. Fixed-point search algorithms have been studied in both the QW [55] and AQC [56] regimes, so interpolation to generate hybrid algorithms should be straightforward. The quantum-walk search on random graphs solved in [31] is based on the same kind of single-avoided-crossing arguments which appear in this work, meaning that these are also natural for hybrid QW-AQC protocols.

Hybrid algorithms such as the ones we presented here can be viewed as particular instances of quantum control techniques applied to solving optimization and search problems. Another application of quantum control to quantum algorithms is based on the Pontryagin minimum principle of optimal control: that optimal control protocols for solving these problems will follow a bang-bang scheme, with successive applications of the extreme values of the controls [57]. An algorithm based on such controls, called the quantum approximate optimization algorithm (QAOA), was proposed by Farhi et al. [58,59]. This protocol can be implemented either through digital quantum circuits or by successively applied Hamiltonians. It has been shown that the QAOA can obtain an optimal \( \sqrt{N} \) scaling in solving the search problem using a transverse field search unitary [60], essentially the problem we consider in this paper.

However, there are two caveats worth noting in terms of the optimality of QAOA-type bang-bang protocols. First, when viewed as an application of successive Hamiltonians, these protocols require infinitely fast switching time, which is generally unphysical. Second, while the optimal control scheme to find the solution is mathematically always of a bang-bang form, this solution may exhibit Fuller’s phenomenon [61,62], in which the optimal solution involves switching back and forth between the two extremal Hamiltonians an infinite number of times in a finite-time window. While mathematically valid, such a control scheme is clearly not physically realizable. It is an open question what happens to a Hamiltonian-based QAOA when finite switching time is added as a constraint. Our result that intermediate protocols between quantum-walk and adiabatic protocols are still able to obtain an optimal speedup provide an encouraging sign
that the QAOA may remain effective with realistic constraints applied.

Recent studies by Muthukrishnan et al. [63,64] on a class of permutation symmetric problems related to, but distinct from, searching have found that, deep in the diabatic regime, the problem can be solved by dynamics which are effectively classical through diabatic cascades. Muthukrishnan et al. focused only on changing the rate of evolution of an AQC algorithm; in contrast, we examined both the shape of the schedule and the rate of evolution. Furthermore, since all of the qubits need to align to interact meaningfully with the energy landscape of the search problem, it is unlikely that a similar classical diabatic cascade regime exists in our study.

In addition to problem size, the performance of a quantum search in a realistic setting will depend on many other factors. By performing a fairly general and multifaceted analysis of such factors, we uncover a landscape where no single protocol dominates. In asymptotically large systems with perfectly specified problems, a straightforward QW approach is best. However, this limit is approached slowly, since the success probability for QW scales only as \( n \), i.e., logarithmically in problem size \( N \). A rich structure exists for computation-ally interesting nonasymptotic sizes. On the other hand, for asymptotically large systems with some degree of problem misspecification, interpolated protocols can outperform the QW approach. A simple open system analysis reveals another layer of structure that can be exploited in realistic settings. Application of these techniques to algorithms with useful applications which can be run on near-future quantum hardware is left to future work [65].

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APPENDIX A: HYPERCUBE OPTIMAL SCHEDULE CALCULATION

Starting from the Hamiltonian for the AQC search on a hypercube (14),

\[ \hat{H}(s) = (1-s) \sum_{j=1}^{n} \frac{1}{2} (1 - \hat{\sigma}_j^{(z)}) + s(1 - |m\rangle \langle m|), \]

we first apply a gauge transformation (a swap of the 1 ↔ 0 labels on a subset of the qubits) to map the marked state \(|m\rangle\) to the state \(|0\rangle\). We then express the Hamiltonian in the symmetric subspace in terms of total spin operators

\[ \hat{S}_a = \frac{1}{2} \sum_{j=1}^{n} \hat{\sigma}_a^{(j)} \quad (A1) \]

for \( a \in \{x, y, z\} \), which have eigenstates \(|\frac{n}{2} - r\rangle_a\) for \( r \in \{0, \ldots, n\} \). In this representation, the marked state is \(|\frac{n}{2}\rangle\); and the AQC search Hamiltonian becomes

\[ \hat{H}(s) = (1-s)(\frac{n}{2} - \hat{S}_z) + s(1 - |\frac{n}{2}\rangle \langle \frac{n}{2}|). \quad (A2) \]

Following Farhi et al. [34], to analyze the eigensystem we obtain the eigenvalue equation

\[ \frac{1 - s}{s} = \frac{1}{N} \sum_{r=0}^{n} \left( \frac{n}{r} \right) \frac{1}{r - \lambda} \quad (A3) \]

for the energy eigenvalues \( E_s = s + (1-s)\lambda \). Farhi et al. [34] solved this at the minimum gap, which occurs at \( s = s_m \) for

\[ \frac{1 - s_m}{s_m} = \frac{1}{N} \sum_{r=0}^{n} \left( \frac{n}{r} \right) \frac{1}{r} - \lambda \equiv R_1, \quad (A4) \]

and showed that \( \lambda_{0,1}^{(\text{gmin})} \approx \pm n/2\sqrt{N} \) for the two lowest eigenvalues corresponding to the ground state \( E_0 \) and first excited state \( E_1 \).

To obtain the optimal schedule following the method in Roland and Cerf [21], we need an expression for the gap as a function of \( s(t) \), not just at the minimum gap. We expand the eigenvalue equation \((A3)\) for \( \lambda \ll 1 \),

\[ \frac{1 - s}{s} = -\frac{1}{N\lambda} + \frac{1}{N} \sum_{r=1}^{n} \left( \frac{n}{r} \right) \frac{1}{r} (1 + \lambda/r) + O(\lambda^2). \quad (A5) \]

Using \( R_1 \) and \( R_2 \) from Eqs. \((10), (23)\), we obtain

\[ \frac{1 - s}{s} = -\frac{1}{N\lambda} + R_1 + \lambda R_2. \quad (A6) \]

This quadratic equation in \( \lambda \) has roots

\[ \lambda = \frac{1}{2R_2} \left\{ \frac{1 - s}{s} - R_1 \right\} \]

\[ \pm \frac{1}{4} \left\{ \frac{1}{R_2} \left( \frac{1 - s}{s} - R_1 \right)^2 + \frac{4}{NR_2} \right\}^{1/2} \quad (A7) \]

and gives, for the gap \( g(s) = (1-s)(\lambda_1 - \lambda_0) \),

\[ g(s) = (1-s) \left\{ \frac{1}{R_2} \left( \frac{1 - s}{s} - R_1 \right)^2 + \frac{4}{NR_2} \right\}^{1/2}. \quad (A8) \]

To optimize the schedule, we need to solve Eq. \((16)\),

\[ \frac{ds}{dt} \leq \epsilon \frac{g^2(s)}{\left| \left. \frac{dH}{ds} \right|_{0,1} \right|}. \]

using the expression for \( g(s) \) in Eq. \((A8)\). To obtain a suitable approximate value for \( \left| \left. \frac{dH}{ds} \right|_{0,1} \right| \), we first calculate \( \frac{dH}{ds} \) in the symmetric subspace representation of Eq. \((A2)\),

\[ \frac{dH}{ds} = -\left( \frac{n}{2} - \hat{S}_z \right) + \left( 1 - \frac{n}{2} \right). \quad (A9) \]
It is sufficient to use the maximum value of \( \frac{dI}{ds} \) \( s = 0 \), which occurs at \( s_m \), where the eigenstates \( |E_{1,0}\rangle = (|\frac{1}{2}\rangle + |\frac{1}{2}\rangle)/\sqrt{2} \), giving \( \frac{dI}{ds} \) \( s = 0 \) \( \leq \frac{\pi}{4} \). We then have the following equation to solve for \( s(t) \):

\[
\frac{ds}{dt} = \frac{4e}{nR_2^2} (1 - s)^2 \left( \frac{1 - s - R_1}{s - R_1} \right)^2 + \frac{2R_2}{N} \]

This can be integrated to obtain

\[
\frac{4et}{nR_2^2} + c = \frac{R_1^2 - 4R_2/N}{2\sqrt{R_2/N}(R_1^2 + 4R_2/N)^2} \times \arctan \left\{ \frac{(1 + R_1)^2 + 4R_2/N}{2\sqrt{R_2/N}} \right\} \left( \frac{1}{1 - s(R_1^2 + 4R_2/N)} \right) + \frac{R_1}{(R_1^2 + 4R_2/N)^2} \ln \left\{ \frac{(1 - s - R_1s)^2 + \frac{4R_2^2s^2}{N}}{(1 - s)^2} \right\},
\]

where \( c \) is the constant of integration. To obtain the constant, set \( s = t = 0 \), giving

\[
c' = \arctan \left\{ \frac{(1 + R_1)/\sqrt{N}}{2\sqrt{R_2}} \right\} + \frac{R_1 + 4R_2/N}{2\sqrt{R_2/N}} \times \frac{1}{R_1^2 - 4R_2/N},
\]

where the factors in front of the arctan term have been rearranged to give a more convenient form for the constant. One can then in principle solve for \( s(t) \). However, the terms on the right-hand side, apart from the arctan, are potentially problematic as \( s \rightarrow 1 \). Given that we started with the approximation \( \lambda \ll 1 \), which occurs at the position of the minimum gap, we cannot necessarily expect that the solution will be valid for \( s \rightarrow 1 \). We first note that only the arctan term on the right-hand side gives a schedule that is valid for all \( 0 \leq s \leq 1 \) and it provides a runtime proportional to \( \sqrt{N} \). If we do not discard these extra terms, we can show that they can be neglected, provided we stop the anneal very slightly before \( s = 1 \), but still well past the minimum gap.

To solve for \( s(t) \) retaining the full expression, invert the arctan to give

\[
s(t) = \frac{2\sqrt{R_2}}{\sqrt{N}(1 + R_1)^2 + 4R_2/N} \times \tan \left\{ \frac{8e\sqrt{R_2R_1}t}{n\sqrt{N}R_2} - c'' \right\} + \frac{1}{1 + R_1},
\]

where \( c'' \) now contains the awkward extra terms

\[
c'' = c' - \frac{1}{1 - s(R_1^2 + 4R_2/N)} \left( \frac{R_1^2 + 4R_2/N}{2\sqrt{R_2/N}} \right) - \frac{R_1}{(1 - s(R_1^2 + 4R_2/N))} \ln \left\{ \frac{(1 - s - R_1s)^2 + \frac{4R_2^2s^2}{N}}{(1 - s)^2} \right\}
\]

The arctan argument is large, so the arctan is close to \( \pi/2 \). We note that the extra terms are small for most values of \( s \) and only become large as \( s \rightarrow 1 \). To check when these terms become \( O(1) \), for the first extra term we solve

\[
\frac{s}{1 - s(R_1^2 + 4R_2/N)} \sqrt{\frac{R_2}{N}} \Rightarrow 1 \quad (A15)
\]

to obtain

\[
s \simeq \frac{1}{1 + 2\sqrt{R_2/N} \Rightarrow 1 + 4/n\sqrt{N}} \quad (A16)
\]

This is well past the minimum gap, which occurs at \( s = 1/(1 + R_1) \simeq 1/(1 + 2/n) \). Applying the same procedure to the second extra term gives, to leading order,

\[
s \simeq 1 - e^{-\sqrt{N}/s}, \quad (A17)
\]

which is even closer to \( s = 1 \) and further from the minimum gap. Since the transition probabilities are only significant close to the minimum gap and hence all the important slowing down of the schedule occurs around the gap, what happens this close to \( s = 1 \) has essentially no effect on the success or runtime of the algorithm.

Dropping the extra terms from the solution provides an expression for \( s(t) \),

\[
s(t) = \frac{2\sqrt{R_2}}{\sqrt{N}(1 + R_1)^2} \times \tan \left\{ \frac{8e\sqrt{R_2R_1}t}{n\sqrt{N}R_2} - c'' \right\} + \frac{1}{1 + R_1},
\]

where we have also dropped terms \( O(1) \) and

\[
c'' = \arctan \left\{ \frac{(1 + R_1)/\sqrt{N}}{2\sqrt{R_2}} \right\}. \quad (A19)
\]

Strictly speaking, this is valid for \( s \ll 1 + 4/n\sqrt{N} \), although in fact it is well behaved right up to and including \( s = 1 \). From this we can obtain the runtime

\[
e_{fr} \simeq \frac{\pi}{4} \sqrt{\frac{N}{4}}, \quad (A20)
\]

where the two arctan terms have each been approximated by \( \pi/2 \), since their arguments are large, \( O(\sqrt{N}) \).

**APPENDIX B: NUMERICAL METHODS**

Our numerical calculations were carried out using the Python programming language (both Python 2.7 and Python 3.5), making considerable use of the Numpy, Scipy, and Matplotlib packages [66–69]. High performance computing resources were not used in this study, although some of the simulations took several days to run on standard desktop workstations. Most of the simulations consisted of
solving the time evolution of the quantum search algorithm by numerically integrating the Schrödinger equation using the appropriate Hamiltonian. This was done by diagonalizing the Hamiltonian and exponentiating it in the diagonal basis, before applying it to the wave function. This process was iterated for time-dependent Hamiltonians, rotating from one instantaneous diagonal basis to the next at small time intervals. For the decoherence studies in Sec. VC, the same process was applied to the density matrix, with dephasing operators also applied along with the unitary time evolution.

For larger systems, we can take advantage of the symmetry in the hypercube to map the dynamics to a search on the line with appropriately weighted edges, as given by Eq. (A2). Provided the initial state is also invariant with respect to this symmetry, the evolution will be restricted to this symmetric subspace. This allows us to perform simulations for much larger numbers of qubits \( n \lesssim 100 \) and hence extract reliable information about the scaling with \( n \) from numerics alone. This provides important checks of the validity of the two-level approximations made to facilitate the analytical calculations.

Optimal AQC schedules \( s^{(n)}(\tau) \) were calculated numerically as solutions of Eq. (20), both to check the analytical solutions for the hypercube and because we can solve numerically with fewer approximations than are required to obtain analytical expressions. Specifically, we calculate the gap \( g(s) \) directly from the Hamiltonian eigensystem, rather than expanding about \( g \) directly from the Hamiltonian eigensystem, rather than expanding about \( g \) for \( \tau = 0 \) and \( \tau = 1 \) and \( s \) to be a list of the corresponding values of \( s(\tau) \), obtained by applying \( F^{-1}(\tau) \) to each element of \( \tau \). Defining \( j(s) \) equal to the number of elements in \( s \) which are strictly less than \( s \), we approximate \( F(s) \) numerically by \( \tilde{F}(s) \), where we replace the integral by a finite sum plus linear interpolation. Writing \( \tilde{s}_j = \frac{1}{2}[s \text{LIST}(j(s)) + s \text{LIST}(j(s) + 1)] \),

\[
\tilde{F}(s) = \sum_{j=1}^{j_{\text{max}}} \frac{s \text{LIST}(j + 1) - s \text{LIST}(j')}{N g^2(s^j)} + \frac{s - s \text{LIST}(j(s))}{N g^2(\tilde{s}_j)}, \tag{B2}
\]

where \( N \) is a normalization factor which is included to ensure that \( \tilde{F}(s = 1) = 1 \). It is straightforward to numerically invert \( \tilde{F}(s) \). This can be accomplished by first finding \( j_{\text{max}}(s) \), the largest value of \( j(s) \) for which \( \tilde{F}(s) < \tau \), and then solving

\[
\tilde{F}(s)|_{j(s)=j_{\text{max}}(s)} = \tau \tag{B3}
\]

for \( s \). Based on this numerical function inversion, we define an iterative method of converging on the solution for \( s^{(n)}(\tau) \).

(i) Set a linearly spaced \( s \text{LIST} \in [0,1] \) and \( \tau \text{LIST} \in [0,1] \) each with the same number of elements.

(ii) Using the values of \( s \) in \( s \text{LIST} \), apply \( F^{-1}(\tau) \) to each corresponding element in \( \tau \text{LIST} \) to generate a new \( s \text{LIST} \)

(iii) Repeat step (ii), with the new \( s \text{LIST} \) as input, until it has converged

The advantage of this iterative method is that, at each iteration, more points in \( s \text{LIST} \) will concentrate in areas where \( 1/g^2 \) is larger, for instance, near the dominant avoided crossing. By using the previously calculated \( s \text{LIST} \) as a mesh in the current iteration, the protocol can continuously improve the quality of the numerical inverse with a fixed number of points in \( s \text{LIST} \).


