Notes on the implementation of the Hidden Shift Algorithm

Sebastián Grijalva

github.com/sebgrijalva

We show a superpolynomial oracle separation between the power of adiabatic quantum computation with no sign problem and the power of classical computation.

1 Introduction

The adiabatic algorithm[1] is a proposed algorithm for quantum optimization. In its simplest form, one considers a quantum Hamiltonian which is a sum of two terms, one being proportional to the objective function of some optimization problem and the other so-called "driving term" being a "transverse field" (some non-commuting additional term). One then adiabatically evolves the Hamiltonian from a large value of the transverse field (where the ground state is easy to prepare) to a small value of the transverse field, where the ground state encodes the desired solution of the optimization problem.

Unfortunately, there is tremendous theoretical evidence that gaps for random instances typically become super-exponentially small [2, 3] so that the time required for adiabatic evolution to remain in the ground state is longer than the time required for even a classical brute force search¹. Other authors have shown exponentially small gaps in simple problems[7] and some explicit simple examples show a super-exponentially small gap[8]. Further, separate from any question about the scaling of the gap, numerical experiments have shown that classical algorithms which simulate the quantum dynamics can perform comparably to a quantum device[9].

Nevertheless, it remains of some interest to ask about the computational power of adiabatic quantum computation. If we consider a Hamiltonian $H = sH_1 + (1-s)H_0$ which is a a linear combination of two arbitrary local Hamiltonians H_0 , H_1 , with some parameter s controlling the dynamics, and require that the gap become only polynomially small so that the adiabatic evolution can be performed in polynomial time, then the problem is completely understood: this model is equivalent to standard quantum computation, and can solve any problem in BQP[10].

However, if we restrict to the case that H has no sign problem in the computational basis then the problem remains open. Here "no sign problem" means that in the given basis, all off-diagonal terms in H are negative; this is sometimes termed "stoquastic". This case of no sign problem includes the adiabatic optimization algorithm discussed at the start if H_1 is equal to an objective function and H_0 is equal to $-\sum_i X_i$ with X_i being the Pauli X matrix on the i-th qubit. It is important to emphasize that the arguments of [2] apply even if the driving term has some sign problem; they depend rather on H_1 being an objective function and H_0 being chosen as some sum of local terms.

We remark that there is a problem of "glued trees" [11] for which an exponential speedup over classical is known using a Hamiltonian with no sign problem that changes slowly in time [12], but this problem is very different from the adiabatic annealing considered here. The gap becomes exponentially small and the evolving quantum state has a large overlap with excited states during the evolution. More

¹While some authors have disputed the perturbative calculation[4], we believe that the general mechanism of localization on the hypercube will still apply for random instances with local driving terms. The short path algorithm exploiting a difference between ℓ_1 and ℓ_2 localization may however be able give a super-Grover speedup[5, 6].

generally, if one allows dynamics in excited states, then it is possible to perform universal quantum computation using Hamiltonians with no sign problem[13]. Thus, Hamiltonians with no sign problem are universal (using excited states) and adiabatic evolution is universal (using Hamiltonians with a sign problem). The question we consider is what happens if we impose both these restrictions: no sign problem and adiabatic evolution in the ground state with a gap that is only polynomially small.

One piece of evidence that it may be hard to simulate this adiabatic evolution classically in general is the existence of topological obstructions to the equilibration of path integral Monte Carlo methods[14]. As explained later, these obstructions help motivate the construction here. Further, these obstructions are perhaps the main reason one should be interested in the question: topological obstructions such as difficulty equilibrating between different winding numbers can have an important effect on practical simulations of quantum systems as is well-studied in the condensed matter community[15], and so it would be useful if there were a general classical method that could overcome all such obstructions.

1.1 Problem Statement and Results

In this paper, we address this question. Our results, in an oracle model defined below, will show a superpolynomial separation between the power of adiabatic computation with no sign problem and the power of classical computation. At the same time, our results give no reason to believe that adiabatic computation with no sign problem is capable of universal quantum computation.

We use a number N to parameterize the problem size (for example, in the example above using qubits, the number of basis states in the computational basis is 2^N), and all references to polynomial scaling will refer to this parameter N. We will assume more generally that the number of computational basis states is $\leq 2^{p(N)}$ for some function p which is polynomially bounded, i.e., the computational basis states can be labelled using polynomial space.

We will define a path of Hamiltonians H_s , for s in some interval to be admissible if it satisfies the following properties (when we refer to a parameter s below, it is always assumed to be in the given interval). First, for all s, H_s must have no sign problem in the computational basis. Second, for all s, H_s must be polynomially sparse, meaning that for every computational basis element $|i\rangle$, there are at most poly(N) basis elements $|j\rangle$ such that $\langle j|H_s|i\rangle$ is nonzero. Third, for all s, for every i,j, $|\langle j|H_s|i\rangle| \leq \text{poly}(N)$. Fourth, for all s, $|\partial_s H_s|| \leq \text{poly}(N)$. Fifth, for all s, H_s has a unique ground state and the spectral gap to the first excited state is $\Omega(1/\text{poly}(N))$. Sixth, the length of the interval is poly(N).

Note that the number of basis states $2^{p(N)}$ and the definition of an admissible path both depend upon many polynomials that we have left unspecified. The particular value of these polynomials is not important; for example, for any such $p(\cdot)$ and N, we can define $N' = \lceil p(N) \rceil$, so that the number of basis states is $\leq 2^{N'}$. Then, if the number of queries needed to solve all instances with some given probability is superpolynomial in N, it is also superpolynomial in N'. Similarly, if that gap is lower bounded by 1/poly(N) for some polynomial, it is lower bounded by 1/N'' for N'' = poly(N) and again the number of queries would be superpolynomial in N''. Still these polynomials should be regarded as fixed in the main result: for some specific choice of polynomials, we show a superpolynomial number of queries. For example, the construction we use gives $p(N) = O(N^2 \log(N)^3)$; it can be tightened somewhat. The reason it is convenient to leave these polynomials unspecified is that it simplifies some accounting later: we will often, given some problem, construct a new problem with a larger number of basis states (increasing p(N) so that it is still polynomial) or different gap; any change in the polynomials from this construction is often not stated explicitly.

The interest in the conditions other than the no sign problem condition is that adiabatic evolution on admissible paths can be efficiently simulated on a quantum computer up to polynomially small error; indeed, evolution under a time-dependent Hamiltonian for a time that is poly(N) will give the desired approximation to adiabatic evolution.

We will say that a path H_s for s in some interval [a,b] satisfies the *endpoint condition* if for both s=a and s=b, the ground state of H_s is some computational basis state, i.e., for some i (possibly different for s=a and s=b), $|i\rangle$ is the ground state of H_s (in a slight abuse of language, we will say

that a vector is "the" ground state of a Hamiltonian when of course the ground state is only defined up to phase). We refer to a as the start of the path and b as the end of the path. So, our interest will be in admissible paths of Hamiltonians which satisfy the endpoint condition because this gives a simple example of Hamiltonians for which it is easy to prepare the ground state of H_a and for which one can measure in the computational basis to determine the ground state of H_b . We say that the path satisfies the condition at one endpoint if some some endpoint (either s = a or s = b), the ground state is a computational basis state.

Also, we can easily concatenate admissible paths which satisfy the endpoint condition: given one such path H_s for $s \in [a, b]$ with $|j\rangle$ being the ground state of H_b and another admissible path H'_s for $s \in [b, c]$ with $|j\rangle$ being the ground state of H'_b , we can concatenate the two paths to get a new admissible path if $H_b = H'_b$. Even if H_b differs from H'_b , it is possible to interpolate between H_b and H'_b by a path which first makes the off-diagonal elements of the Hamiltonian tend to zero, then changes the diagonal entries until they agree with diagonal entries of H'_b , then increases the off-diagonal elements until they agree with H'_b ; doing this in an obvious way (for example, linear interpolation) still gives an admissible path from H_a to H'_c .

One might be slightly surprised in one respect at our endpoint condition, though, since in adiabatic evolution with a transverse field, at s=0 the ground state of the Hamiltonian is a uniform superposition of all computational basis states, which may be written as $|+\rangle^{\otimes N}$. However, for a system of N qubits there is an obvious admissible path $H_s=-(1-s)\sum_i Z_i-s\sum_i (X_i)$ for $s\in[0,1]$ with the ground state of H_0 being a computational basis state and the ground state of H_1 being $|+\rangle^{\otimes N}$. So, given some admissible path which satisfies the endpoint condition at the right endpoint and with the Hamiltonian at the left endpoint being $-\sum_i X_i$ (for example, interpolation between a transverse field Hamiltonian and some objective function for an optimization problem), we can concatenate with the path above to get an admissible path which satisfies the endpoint condition.

We will consider a version of the problem with an oracle in order to give a superpolynomial lower bound on the ability of classical algorithms to solve this problem. Our oracle for a Hamiltonian will be similar to those considered previously [16, 17, 18]. The Hamiltonians that we consider can be simulated efficiently on a quantum computer with quantum queries of the oracle [16].

We define problem AdNSP as follows. An instance of this problem is defined by some admissible path H_s which satisfies the endpoint condition. For definiteness, we assume that s lies in the interval [0,1]. A query of the oracle consists of giving it any i which labels some computational basis state $|i\rangle$ as well as giving it any $s \in [0,1]$, and the oracle will return the set of j such that $\langle j|H_s|i\rangle$ is nonzero as well as returning the matrix elements $\langle j|H_s|i\rangle$ for those j to precision $\exp(-\text{poly}(N))$, i.e., returning the matrix elements to poly(N) bits accuracy for any desired polynomial. We will call those j the neighbors of i, and we will say that we query state i at the given s. The oracle will also return the diagonal matrix element $\langle i|H_s|i\rangle$. The problem is: given query access to the oracle, and given the computational basis state $|i\rangle$ which is the ground state of H_0 , determine the computational basis state $|j\rangle$ which is the ground state of H_1 . We say a classical algorithm solves this problem for an instance with some given probability if it returns the correct j with at least that probability. (As remarked above, the definition of an admissible path implicitly depends on various polynomials; so implicitly the problem AdNSP also depends on various polynomials.)

Note that given an unlimited number of queries to the oracle, it is possible to simulate the quantum evolution on a classical computer since one can determine the Hamiltonian to exponentially small error.

Remark: we have stated above that the oracle returns the matrix elements only to polynomially many bits. This restriction is unnecessary for all the lower bounds on queries later, which would still hold even if the oracle returned the matrix elements to infinite precision.

Our main result is:

Theorem 1. For some constant c, for some specific choice of polynomials $p(\cdot)$ and choice of polynomials defining an admissible path, there is no algorithm that solves every instance of AdNSP with probability greater than $\exp(-cN)$ using fewer than $\exp(\Theta(\log(N)^2))$ classical queries.

Throughout, when we refer to an algorithm, the algorithm may be randomized and may take

an arbitrary amount of time. Remark: as is standard terminology, we refer to functions which are $O(\exp(\log(N)^{\alpha}))$ for some fixed α as quasi-polynomial functions, and denote an arbitrary such function by $\operatorname{qpoly}(N)$. A function which is $\exp(\Theta(\log(N)^2))$ is quasi-polynomial but is superpolynomial.

1.2 Outline and Motivation for Proof

The motivation for the proof of Theorem 1 is, to some extent, an idea from [14]: path integral Monte Carlo (which is a very natural classical algorithm for simulating quantum systems with no sign problem) in many cases cannot distinguish between the dynamics of a quantum particle on some graph G and the dynamics on its universal cover \tilde{G} . However, at the same time, the largest eigenvalue of the adjacency matrix of G (which will give us, up to a minus sign, the ground state energy of a Hamiltonian we define for that graph) may be much larger than it is on \tilde{G} (to be precise, for an infinite graph we should not talk about "the largest eigenvalue", but rather use spectral norm), so that a quantum algorithm can distinguish them. We emphasize that if \tilde{G} is a *finite cover* of G, the largest eigenvalue of the adjacency matrix of \tilde{G} is the same as that of G.

This difference on infinite graphs has a finitary analogue: there is a difference between the ground state energy on a complete graph and the ground state energy on a tree graph with the same degree as the complete graph, with the difference in energy persisting no matter how deep the tree is, assuming the degree is ≥ 3 . Here, the ground state energy of a graph is minus the largest eigenvalue of the adjacency matrix of that graph.

Our proof is based on the following main idea: we construct two different graphs which have different ground state energies, but for which we can give a superpolynomial lower bound on the number of classical queries to distinguish those graphs; we quantify this ability to distinguish the graphs in terms of mutual information between a random variable which is a random choice of the graphs and another random variable which is the query responses. We will term these graphs C and D; these actually refer to families of graphs depending on some parameters.

The proof has two main parts: first, using these graphs to construct a family of instances of AdNSP which cannot efficiently be solved with classical queries, and, second, proving the lower bound on the number of classical queries. Proving spectral properties of the quantum Hamiltonians of these graphs is an additional part of the proof, but is relatively simple.

The first part of the proof is in Section 2, where we show that it suffices to prove Theorem 1 in a different query model. This part of the proof is perhaps less interesting than later parts of the proof, though it is important to understand the modified query model that we define. In this query model, the oracle gives less information in response to queries, making it impossible to distinguish between a graph G and some cover \tilde{G} . Each vertex of the graph corresponds to a computational basis state so that neighbors of a vertex are also neighbors that one might receive in response to a query. On G, one might follow some cycle on the graph, returning to the start, but if \tilde{G} is the universal cover this is not possible: in the modified query model, one will not be able to know that one has returned to the start if one follows a cycle.

Then, in Section 3, we then reduce the problem of proving Theorem 1 to showing two graphs C, D satisfying certain properties exist. The needed properties of the graphs are summarized briefly in Table 1. The main result is Lemma 3.

The rest of the paper is concerned with constructing C, D. In Section 4 we give a first attempt at a construction, taking C to be a complete graph of O(1) vertices and D to be a bounded depth tree graph (an infinite tree graph would give an example of a cover of C). The idea is that if one does not reach a leaf of D, then D is indistinguishable from the cover of C, and by choosing the height of the tree superpolynomially large, it may take superpolynomially many queries to reach a leaf.

Unfortunately, the construction of Section 4 suffers from two serious defects. The first is that the it uses "too many" states, i.e., number of vertices in the graph is not $\exp(\operatorname{poly}(N))$ if we take the height of the tree superpolynomially large. The second and more serious defect is that the graphs come with a privileged vertex called the "start vertex", and in the modified query model we will still be able to determine when we return to the start vertex; a random walk in C will often return to the start

vertex but in D one will not so a classical algorithm can efficiently distinguish them. Nevertheless, this construction is worthwhile as it introduces certain key ideas used later.

The second part of the proof, constructing C and D fulfilling all needed properties (including that they cannot be efficiently distinguished by any classical algorithm and that there are only $\exp(\operatorname{poly}(N))$ vertices in the graph), starts in Section 5. Here, we introduce the notion of a "decorated graph". The idea is to define some sequence of tree graphs for which it is "hard" in some sense to reach certain vertices far from the start vertex because one tends (speaking very heuristically) to get "lost" in other paths near the root. This tendency to get lost will make it hard for classical algorithms to detect the difference between C and D.

In that section, we also give bounds on the energy of the Hamiltonians corresponding to these graphs and prove some properties of the ground states.

In Section 6, we give lower bounds on the number of classical queries needed to distinguish between C and D. Lemma 5 quantifies the difficulty of distinguishing C and D in terms of mutual information; Lemma 5 takes as input an assumption about difficulty of "reaching" a certain set of vertices Δ in D using queries starting from a given "start vertex". Difficulty of reaching this set follows from an inductive Lemma 6.

The results in Section 5 and Section 6 are given in terms of a number of parameters. In Section 7 we fix values for these parameters and prove Theorem 1.

In Section 8 we briefly discuss a case of linear interpolation rather than arbitrary paths.

2 Modifications to Query Model

This section consists of two different subsections, which allow us to consider a more restrictive query model that we call the modified query model.

2.1 Related States

We first show that we may, in everything that follows, assume that every state queried is either the initial state $|i\rangle$ which is the ground state of H_0 or a neighbor of some state queried in a previous query. For example, it may query i for some value of s, receiving neighbors j_1, j_2, \ldots . It may then choose to query j_1 (possibly for some different s), receiving neighbors k_1, k_2, \ldots , at which point it may query any of $i, j_2, j_3, \ldots, k_1, k_2, \ldots$, but it will never query an "unrelated state", meaning a state (other than $|i\rangle$) that it has not received in response to a previous query.

This result is essentially the same as Lemma 4 of the arXiv version of [11]. The basic idea of the proof is one that we will re-use in Section 2.2. At a high level, the idea is: given any problem H_s from an instance of AdNSP, we will construct some new oracle which is "weaker" in some sense than the original oracle; in this case, whenever a related state is queried it returns the same responses as the original oracle, but whenever an unrelated state is queried, it returns some fixed response (i.e., the same response for any H_s) which hence gives no information about H_s . Thus, queries of the weaker oracle can be simulated by queries of the original oracle (simply replace the response to a query of an unrelated state by this fixed response) but not vice versa. We then construct (for any H_s) some set of paths H'_s which corresponds to some other instances of AdNSP such that the original oracle for H'_s is almost equivalent to the weaker oracle for H_s . Here, "almost equivalent" means that for a random choice of path H'_s from this set, with probability close to 1 the original oracle returns the same responses as does the weaker oracle for H_s . Finally, since by assumption A solves every instance of AdNSP with probability p, including in particular the paths H'_s , we can define algorithm A' to be given by algorithm A using queries of the weaker oracle for H_s . This construction will imply some change in the polynomials defining AdNSP; in particular, in this case the size of the Hilbert space will change.

Formally:

Lemma 1. For any algorithm A that solves every instance of AdNSP with probability $\geq p$ using only quasi-polynomially many queries and possibly using queries of unrelated states, there is some algorithm A' which only queries related states and solves every instance of AdNSP with probability $\geq p - \text{qpoly}(N)2^{-N}$ using at most as many queries as A (albeit with some change in the polynomials defining AdNSP). Hence, if algorithm A succeeds with probability large compared to $\text{qpoly}(N)2^{-N}$, then algorithm A' succeeds with a probability that is comparable to that of A.

Proof. Given any path H_s with Hilbert space \mathcal{H} of dimension $\dim(\mathcal{H}) = 2^{p(N)}$, consider a new Hilbert space \mathcal{H}' of dimension $\dim(\mathcal{H}') = 2^{p(N)+N}$ which is exponentially larger. Define a path H_s' by

$$H_s' = \Pi \begin{pmatrix} H_s & \\ & WI \end{pmatrix} \Pi^{-1}, \tag{1}$$

where the rows and columns correspond to computational basis states, where the first block is of size $2^{p(N)}$ and the second block is of size $2^{p(N)+N}-2^{p(N)}$, where Π is a permutation matrix chosen uniformly at random, and where I is the identity matrix and W is a scalar. We choose W larger than the largest eigenvalue of H so that the ground state of $\begin{pmatrix} H_s \\ WI \end{pmatrix}$ is completely supported in the first block and is given in the obvious way from the ground state of H_s , and hence the ground state of H_s is given by applying Π to that state.

Suppose on the q-th query, the algorithm queries an unrelated state. The number of unrelated states whose image, under Π^{-1} , is in the first block is at least $\dim(\mathcal{H}') - \dim(\mathcal{H}) - q$. Since the permutation Π is random, with probability at least $(\dim(\mathcal{H}') - \dim(\mathcal{H}) - q)/\dim(\mathcal{H}')$ the response to the query will be that the state has no neighbors and that the diagonal matrix element of that state is W. Call this "response R".

If an algorithm makes only quasi-polynomially many queries, with probability $\geq 1 - \operatorname{qpoly}(N) 2^{-N}$ the response to all queries of unrelated states will be R. So, given some algorithm A which may query unrelated states, and which makes only quasi-polynomially many queries, we may define a new algorithm A' which modifies A by assuming (without querying the oracle) that the response to any query of an unrelated state will be R. Remark: if A' finds some inconsistency in this assumption, for example if it queries some unrelated state and assumes the response is R and then later that state is returned as the neighbor of some previous query, algorithm A' will terminate and return some arbitrary result.

Then, A' queries only related states and, with probability $\geq 1 - \operatorname{qpoly}(N) 2^{-N}$, algorithm A returns the same result as algorithm A' does. Here "probability" refers to both random choice of Π and randomness in A; if A is randomized, of course we assume that A' uses the same source of randomness. Each instance of H'_s is defined by an instance of H_s and by a choice of Π . If A returns the correct result for every instance of H'_s with at least probability p for some p, then, trivially, for any H_s the average over Π of its probability of returning the correct result is at least p. Hence, for any H_s , the probability that A' returns the correct result is at least $p - \operatorname{qpoly}(N) 2^{-N}$.

2.2 Modified Query Model

We now introduce the modified query model in contrast to the query model given previously (which we will refer to as the original query model). We show that if Theorem 1 holds using the modified query model, then it holds using the original query model. Very briefly: the modified query model will be such that if the algorithm follows some nonbacktracking path of queries that forms a cycle (for example, querying i to get some neighbor j, querying j to get some neighbor k, querying k to get i which is a neighbor of k), then the query responses will make it impossible to determine that one has returned to the start of the cycle (in this case, i).

To explain the modified query model in more detail, we have an infinite set of *labels*. Each label will correspond to some computational basis state, but the correspondence is many-to-one; we describe this correspondence by some function $F(\cdot)$. The algorithm will initially be given some label l that

corresponds to the computational basis state $|i\rangle$ that is the ground state of H_0 . A query of the oracle consists of giving it any label m that is either l or is a label that the algorithm has received in response to some previous query, as well as giving it any $s \in [0,1]$, and the oracle will return some set S of labels such that F(S) is the set of j such that $\langle j|H_s|F(m)\rangle$ is nonzero. Distinct labels in S will have different images under $F(\cdot)$ so that |S| is equal to the number of neighbors.

The oracle will also return, for each label $n \in S$, the matrix elements $\langle F(n)|H_s|F(m)\rangle$ to precision $\exp(-\text{poly}(N))$, i.e., returning the matrix elements to poly(N) bits accuracy for any desired polynomial. The oracle will also return the diagonal matrix element $\langle F(m)|H_s|F(m)\rangle$. The labels in S will be chosen as follows: if label m was received in response to some previous query on a label n, so that $\langle F(n)|H_s|F(m)\rangle$ is nonzero and hence $F(n) \in F(S)$, then label n will be in S, i.e., we will "continue to label F(n) by label n". However, for all other j such that $\langle j|H_s|F(m)\rangle$ is nonzero, we will choose a new label (distinct from all previous labels) to label the given vertex j, i.e., a new label o such that F(o) = j.

Thus, after a sequence of queries by the algorithm, we can describe the queries by a tree, each vertex of which is some label, with neighboring vertices in the tree corresponding to computational basis states which are neighbors.

We use the same idea as in the proof of Lemma 1. In this case, the weaker oracle is the oracle of the modified query model. This can clearly be simulated by the original oracle, since one can simply invent new labels for a state if the oracle gives one a label that one has seen previously, but not necessarily vice versa.

We define, for each Hamiltonian H, a model which has a large but finite set of labels. The function $F(\cdot)$ mapping labels to vertices will be 2^N -to-one. If Hamiltonian H acts on Hilbert space \mathcal{H} , then these labels l will correspond one-to-one to computational basis states of some Hilbert space \mathcal{H}' with dimension $2^N \dim(\mathcal{H})$. We will define a Hamiltonian H' acting on \mathcal{H} as follows.

Label computational basis states of \mathcal{H}' by a pair i, x where i is a computational basis state of \mathcal{H} and x is a bit string of length N. For each $a \in \{0, 1, ..., N-1\}$, let X_a denote the operator that flips the a-th bit of this bit string, i.e.,

$$X_a = \sum_{i,x} |i, x \oplus 1_a\rangle\langle i, x|,$$

where 1_a is a binary vector with entry 1 in the a-th position, and 0 elsewhere, and \oplus is the exclusive OR operator. Thus, one may regard that N bits of the bit string as additional qubits and X_a as the Pauli X operator on them.

For each pair i, j of computational basis states of \mathcal{H} , choose randomly some permutation $\pi_{i,j}(\cdot)$ of the bit strings of length N. Here we emphasize that this is *not* a permutation of the individual bits of the string, but rather a permutation from a set of size 2^N to itself. Choose these permutations uniformly and independently subject to the condition that $\pi_{j,i}$ is the inverse function of $\pi_{i,j}$. Define H' by

$$H' = -T \sum_{a} X_a - \sum_{i,j,x} |i,x\rangle\langle j,\pi_{i,j}(x)| \Big(\langle i|H|j\rangle\Big), \tag{2}$$

where T > 0 is a scalar and the second term, in words, means that for each pair i, j of computational basis states of \mathcal{H} , for each x, if there is a matrix element of H between i and j, then there is a matrix element of H' between i, x and $j, \pi(x)$.

Remark: we have written H' using a particular choice of basis states. However, we can, as in Eq. (1), assume that H' is conjugated by a further random permutation so that the algorithm has no information on the labels of the basis states. In this case, if the algorithm receives some state $|i,x\rangle$ as a neighbor in response to some query, and some other state $|i,y\rangle$ in response to a query with $x \neq y$, the algorithm will be unable to know that in both cases the first index i is the same.

The key idea of this construction, using this remark, is that we can ensure that the algorithm is exponentially unlikely to receive the same label twice in response to a query, except for some trivial situations.

Given a path of Hamiltonians H_s , we can define a path H'_s in the obvious way. Define an isometry \mathcal{L} from \mathcal{H} to \mathcal{H}' by

$$\mathcal{L} = 2^{-N/2} \sum_{i,x} |i,x\rangle\langle i|.$$

Choosing T > 0, the ground state subspace of $-T \sum_a X_a$ is the range of \mathcal{L} , and we may choose T large but still T = O(poly(N)) so that the ground state of H'_s is given, up to polynomially small error by

$$\psi_s' = \mathcal{L}\psi_s$$

where ψ_s is the ground state of H_s ; the error can be bounded by any desired polynomial by choosing T to be a sufficiently large polynomial function of N. Further, if the path H_s is admissible, so is the path H_s' ; in particular, we can choose T large enough but still O(poly(N)) such that the fifth condition on spectral gap is satisfied.

The path H'_s does not yet satisfy the endpoint condition. However, this is easy to resolve. Concatenate the path H'_s with a final path along which the term $-T\sum_a X_a$ is replaced by the more general term

$$-T\left(\cos(\theta)\sum_{a}X_{a}+\sin(\theta)\sum_{a}Z_{a}\right),$$

with Z_a being the Pauli Z operator on the additional qubits, i.e., $Z_a = \sum_{i,x} (1 - 2x_a)|i,x\rangle\langle i,x|$, where x_a is the a-th entry of bit string x. On this final path, vary from $\theta = 0$ to $\theta = \pi/2$. Similarly, also concatenate with an initial path along which we vary from $\theta = \pi/2$ to $\theta = 0$. Let \tilde{H}_s denote the path of Hamiltonians given by H'_s concatenated with these two additional paths. Then, if the ground state of H_0 is $|i\rangle$, the ground state of the \tilde{H}_s at the start of the path is given by $|i,0\rangle$, and similarly at the end of the paths.

Thus, we have an admissible path of Hamiltonians \tilde{H}_s satisfying the endpoint condition, with the ground state of \tilde{H}_s trivially related to that of H_s for s=0,1 (the polynomially small error in $\psi'_s=\mathcal{L}\psi_s$ at intermediate steps of the path is unimportant for this).

Suppose now we give the algorithm some additional information in response to queries: if we query some state $|i,x\rangle$ and a neighbor is some other state $|i,y\rangle$, then the algorithm will be informed that the value of i remains the same. This additional information can only help. However, we claim that with this additional information, up to exponentially small error, the queries of \tilde{H}_s in the original query model can be described by queries of H_s in the modified query model. More precisely, assume we know that \tilde{H}_s is given by this construction. Then, the only information given by querying \tilde{H}_s along the "final" or "initial" paths above where θ varies is that one may get multiple labels which are known to have the same first index, i.e., one may start with $|i,0\rangle$, and then get labels describing other states $|i,x\rangle$. Further, we claim that up to exponentially small error explained below, queries along the path H'_s in the original query model can be described by queries of H_s in the modified query model.

To show this, consider the probability that the algorithm receives the same label twice in response to a query. Suppose the algorithm makes multiple queries in which the first label does not change and the algorithm knows it due to the additional information above. Thus, the algorithm will know that some set of labels will describe the same value of the first index. After some number of queries, there will be several sets S_1, S_2, \ldots where each set is a set of labels known to describe the same value of the first index. Formally, there is an equivalence relation on labels: two labels are equivalent if one label is received in response to a query on the other and it is known that the first index did not change, and we extend this equivalence transitively, and the sets S_1, S_2, \ldots are equivalence classes under this relation. Now, consider the probability that some query of some label $|i,x\rangle$ gives a label $|j,y\rangle$ that has been seen previously by the algorithm in response to a previous query, in the case that $j \neq i$ (so that this query does not simply increase the size of one of the equivalence classes, but actually yields new information about the Hamiltonian). The second index y equals $\pi_{j,i}(x)$ and $\pi_{j,i}(x)$ is chosen uniformly at random subject to the condition that $\pi_{j,i}$ is the inverse of $\pi_{i,j}$. Hence, after only quasi-polynomially many queries (so that only quasi-polynomially values of $\pi_{j,i}$ have been fixed) it is exponentially unlikely that $\pi_{j,i}(x)$ will agree with any previously given value of the second index, unless it is the case that we

have previously queried $|j, \pi_{j,i}(x)\rangle$, i.e., unless $|i, x\rangle$ was received as a label of a neighbor of $|j, \pi_{j,i}(x)\rangle$, which is precisely the case in the modified query model that we receive the same label for a given value of the state. Now consider a query in which the first index does not change; suppose we queried a vertex in some equivalence class S, receiving some new label $|i, x\rangle$. It is exponentially unlikely that this label labels a state in some other equivalence class, though it may be only polynomially unlikely that it labels a state in the given class S. Hence, except for an exponentially small probability, a query in which the first index does not change will not collapse two different equivalence classes.

Hence we have:

Lemma 2. If Theorem 1 holds in the modified query model, then it holds in the original query model.

3 Distinguishing Graphs

Now, within the modified query model we show how to use two (families of) graphs C, D to construct instances of AdNSP to prove Theorem 1. The main result is Lemma 3, which we give at the end of this section after developing the machinery of paths needed. In this section we assume that several properties of C, D hold. We summarize these in Table 1. More detail is given below and these properties are proven in later sections of the paper

Both graphs C, D will have a privileged vertex called the "start vertex". For tree graphs, the start vertex will often be the root of the tree.

Given a graph G, we say that the Hamiltonian corresponding to that graph is equal to minus the adjacency matrix of that graph, where each vertex of the graph corresponds to a distinct computational basis state. We will assume that the ground state energy of the Hamiltonian of C is lower than the ground state energy of D by a spectral gap that is $\Omega(1/\text{poly}(N))$; indeed, the difference will be much larger than that in our construction. We will also assume that the gap between the ground state of the Hamiltonian of C and the first excited state of that Hamiltonian is also $\Omega(1/\text{poly}(N))$; indeed, that difference is also much larger than that. Further, we will assume that the amplitude of the ground state wavefunction of C on the start vertex is also $\Omega(1/\text{poly}(N))$.

At the same time, we will also assume a superpolynomial lower bound on the number q of classical queries needed to distinguish C from D in the modified query model above, assuming that the first vertex queried is the start vertex. The modified query model refers to querying a Hamiltonian; here the Hamiltonian will be the Hamiltonian of the given graph, so that computational basis states are neighbors if the corresponding vertices are neighbors. The bound is given in terms of mutual information in Lemma 5; if the algorithm is randomized, then the mutual information is conditioned on any randomness used by the algorithm. Later choices of constants in Section 7 will make $q = \exp(\Theta(\log(N)^2))$.

- 1. The ground state energy of the Hamiltonian of C lower than that of D by $\Omega(1/\text{poly}(N))$.
- 2. $\Omega(1/\text{poly}(N))$ gap of Hamiltonian of C.
- 3. Amplitude of ground state of C on start vertex is $\Omega(1/\text{poly}(N))$.
- 4. Lower bound on number of classical queries to distinguish C from D in the modified query model. Precisely: if the graph is chosen randomly to be C with probability 1/2 and D with probability 1/2, then with fewer than q classical queries the mutual information (in bits) between the query responses and the choice of graph is bounded for all sufficiently large N by some quantity which is strictly less than 1, for some q which is superpolynomial.
- 5. Number of vertices is $O(2^{\text{poly}(N)})$. This property is needed because each vertex will correspond to some computational basis state.

Table 1: List of properties needed for graphs ${\cal C}$ and ${\cal D}.$

Then, given these graphs, we now construct a path of Hamiltonians H_s . To describe this path, we start with a simplified case. Consider a problem with computational basis states labelled by vertices of a graph G (where G is either C or D) as well as by some additional basis state $|0\rangle$. We will label the basis state corresponding to the start vertex of G by $|s\rangle$ (hopefully no confusion will arise with the use of s as a parameter in the path). Consider the two parameter family of Hamiltonians

$$H(t,U) = -U|0\rangle\langle 0 + t(|0\rangle\langle s| + h.c.) + H(G),$$
(3)

where H(G) is the Hamiltonian corresponding to the graph G. We take t < 0 so that the Hamiltonian has no sign problem.

Now consider a path of Hamiltonians starting at very negative U and with t=0 (so that initially the ground state is $|0\rangle$ for both C and D), then increasing t slightly and increasing U, followed by decreasing U. At the end of this path, if G=D, the ground state of the Hamiltonian will still be $|0\rangle$ but if G=C, the ground state of the Hamiltonian will be the ground state of H(C).

We now show that this is possible with an admissible path assuming the properties of C and D above. For both C and D, the ground state energy of H(G) is only $\operatorname{poly}(N)$ so we may take U only $\operatorname{polynomially}$ large initially. Giving the rest of the path in detail, increase t to an amount $\Omega(1/\operatorname{poly}(N))$ and then change U so that it is $1/\operatorname{poly}(N)$ larger than the ground state energy of H(C), but still much smaller than the ground state energy of H(C). Do this with t much smaller than the spectral gap of H(C) and much smaller than the difference between the ground state energy of H(C) and the ground state energy of H(C); given the differences in ground state energies of H(C) and H(D) and the gap of H(D), it is possible to do this with t that is indeed $\Omega(1/\operatorname{poly}(N))$ so that the gap of the Hamiltonian H(t,U) remains $\Omega(1/\operatorname{poly}(N))$. Finally decrease t to 0.

For use later, let us call the path defined in the above paragraph P(G).

At first sight, this path P(G), combined with the lower bound 4 of Table 1 might seem to solve the problem of the needed separation between problems in AdNSP and the power of classical algorithms: the classical algorithm cannot distinguish the two graphs but one can distinguish them with an admissible path of Hamiltonians. However, this is not true; for one, our path of Hamiltonians does not satisfy the endpoint condition as the ground state of the Hamiltonian at the end of the path is a superposition of basis states. Further, the problem is to compute the basis vector at the end of the path, not to distinguish two graphs².

We might try to solve this by concatenating the path of Hamiltonians above with an additional path that decreases H(G) to zero while adding a term $V|s\rangle\langle s|$ and with V decreasing from zero, so that the ground state of the Hamiltonian (in the case that G=C) evolves from being the ground state of H(G) to being $|s\rangle$ and the path now satisfies the endpoint condition.

This still however does not solve the problem: for this path H_s , a classical algorithm can determine the ground state at the end of the path (which we will assume to occur at s=1) by querying the oracle three time, first querying $\langle 0|H_1|0\rangle$, then querying the oracle to find the neighbors of $|0\rangle$ in the middle of the path (so that it can determine s), and finally querying $\langle s|H_1|s\rangle$.

So, to construct the path showing Theorem 1, we use an additional trick. First, we consider N different copies of the problem defined by Hamiltonian Eq. (3) "in parallel". Here, taking "copies in parallel" means the following, given several Hamiltonians H_1, H_2, \ldots, H_N , with associated Hilbert spaces $\mathcal{H}_1, \mathcal{H}_2, \ldots$, we define a Hamiltonian H on Hilbert space $\mathcal{H} \equiv \mathcal{H}_1 \otimes \mathcal{H}_2 \otimes \ldots$ by $H = H_1 \otimes I \otimes I \ldots + I \otimes H_2 \otimes I \otimes \ldots + I \otimes H_2 \otimes I \otimes \ldots + I \otimes I$ is the identity matrix. There is an obvious choice of computational

²Remark: we could have defined a different oracle problem which differs from AdNSP in two ways: first, we only require that the admissible path satisfy the endpoint condition at endpoint s=0 and second, we say that an algorithm "solves" the problem if it computes the amplitude of the basis state $|i\rangle$ which is the ground state of H_0 in the wavefunction which is the ground state of H_1 to within error 1/poly(N) with success probability $\geq 2/3$. Then, for graph C this amplitude is 0 and for D this amplitude is 1 but the classical algorithm cannot distinguish the two cases with probability much larger than 1/2. If we considered this class, then the given path would prove the needed separation. However, this is not what we are considering.

basis states for \mathcal{H} , given by tensor products of computational basis states for $\mathcal{H}_1, \mathcal{H}_2, \ldots$ Similarly, given paths of Hamiltonians, we consider the paths in parallel in the obvious way. If each path is admissible, then the path given by those paths in parallel is also admissible; note that the size of the Hilbert \mathcal{H} is $2^{\text{poly}(N)}$ if each \mathcal{H}_i has dimension $2^{\text{poly}(N)}$.

For each of these N copies, we choose G to be either C or D independently, so that there are 2^N possible instances. Write G_i to denote the graph chosen on the i-th copy. For each copy i let P_i denote the path P for that copy. Let \tilde{P} denote the path given by taking all those path P_i in parallel. At the end of the path \tilde{P} , the ground state is a tensor product of $|0\rangle$ on some copies and the ground state of H(C) on some other copies.

To give an intuitive explanation of the trick we use, it will be that we use this property of the ground state as a kind of key to find an entry in a database: there will be some projector (Π below) which is diagonal in the computational basis and one must find entries of it which are nonzero, and the adiabatic evolution will use this property of the ground state to find it, but the classical will not be able to.

The trick is: concatenate that path \tilde{P} with a further path Q. To define this path Q, add two additional terms to the Hamiltonian

$$V\sum_{i}|s\rangle_{i}\langle s|+W\Pi,$$

where $\sum_i |s\rangle_i \langle s|$ denotes the projector onto $|s\rangle$ on the *i*-th copy tensored with the identity on the other copies, where V,W are scalars, and where Π is a projector which is diagonal in the computational basis. The projector Π will be equal to 1 on a given computational basis state if and only if that computational basis state is a tensor product of basis state $|0\rangle_i$ on all copies for which $G_i = D$ and of basis states corresponding to vertices of G_i for all copies on which $G_i = C$. Thus, the ground state at the end of path \tilde{P} is an eigenvector of P with eigenvalue 1. The path Q is then to first decrease W from zero so that it is large and negative (it suffices to take it polynomially large) while keeping t = 0; then decrease the coefficient in front of H(G) to zero while increasing V to be $\Omega(1/\text{poly}(N))$.

Making W large and negative ensures that throughout Q, the ground state of the Hamiltonian is in the eigenpace of Π with unit eigenvalue. This decrease in the coefficient in from of H(G) combined with increase in V ensures that the ground state at the end of the path is a computational basis state: it is a tensor product of $|0\rangle_i$ on all copies for which $G_i = D$ and of states $|s\rangle_i$ for all copies on which $G_i = C$. We choose V to be $\Omega(1/\text{poly}(N))$ so that the gap of the Hamiltonian will be $\Omega(1/\text{poly}(N))$.

Let \hat{P} be the concatenation of \tilde{P} and Q. Note that there are 2^N possible instances of path \hat{P} , depending on different choices of G_i .

Now we bound the probability of a classical algorithm to determine the final basis state. This lemma shows that if we construct graphs which satisfy **1-5** of Table 1, then Theorem 1 follows.

Lemma 3. If items **1-3,5** of Table 1 hold, then \hat{P} is an admissible path. Further, no algorithm which uses a number of queries which is quasi-polynomial and is smaller than q can solve all instances with probability greater than $\exp(-cN)$.

Proof. By construction \hat{P} is admissible.

Choose each G_i independently, choosing it to be C with probability 1/2 and D with probability 1/2. Suppose that with fewer than q queries, the mutual information between G_i and the query responses is bounded by S < 1. Hence, the average entropy of G_i given the query responses is at least 1 - S so, since the entropy of G_i given the query responses is bounded by 1, with probability at least (1 - S)/2/(1 - (1 - S)/2) the entropy of G_i given the query responses is at least (1 - S)/2.

Thus, with probability that is $\Omega(1)$, the entropy of G_i is $\Omega(1)$. To get oriented, assume that these events (the entropy of each G_i) are independent; that is, define N additional binary random variables S_i which quantify the entropy of G_i being $\geq (1-S)/2$ or not, and assume that these are all independent. Then, with probability $1 - \exp(-\Omega(N))$, there are $\Theta(N)$ independent variables G_i each of which have entropy $\Omega(1)$ and so it is not possible to determine all G_i with probability better than $\exp(-cN)$ using only quasi-polynomially many queries.

Finally, consider the possibility that the S_i are not independent. For example, there is a rather silly algorithm that makes these S_i dependent on each other: consider any algorithm A that gives independent S_i and define a new algorithm A' that calls A with probability 1/2 and makes no queries with probability 1/2 (in which case all $S_i = 1$ since no information is known about any G_i). Then the variables S_i for A' are not independent. However, this "silly algorithm" certainly does not help.

Still we must consider the possibility that there is some way of correlating the S_i which would help. Suppose there were an algorithm which gave correlated S_i , so that for some i the mutual information between G_i and query responses, conditioned on some responses for $j \neq i$ and conditioned on the G_j for $j \neq i$, was larger than S. However, we could then postselect this algorithm on the query responses to the set of $j \neq i$ to give an algorithm that just acted on copy i but which gave mutual information greater than S.

4 A First Attempt

In this section we give a first attempt at constructing two graphs, C and D which satisfy the properties of Table 1. Unfortunately, the example will not quite satisfy the fourth property (using the privileged start vertex it will be possible to efficiently distinguish them, but it will not be possible without that knowledge) or for the fifth: the graph D will have too many vertices. The construction later will fix both of these defects.

Briefly, the graph C is a complete graph on 4 vertices, i.e., every one of the four vertices has degree 3 so it connects to every other vertex. The graph D is a tree graph where every vertex except the leaves has degree 3, i.e., D is given by attaching three binary trees to some root vertex. We choose all the leaves of D to be at distance h from the root for some h so that D has $1+3+2\cdot 3+2^2\cdot 3+\ldots+2^{h-1}\cdot 3$ vertices.

We choose any vertex of C arbitrarily to be the start vertex. We choose the start vertex of D to be the root. Then, it is trivial to verify item 1-3 of Table 1.

Consider what it means to distinguish two graphs in the modified query model. A query of any vertex can return only the information of the degree of that vertex and whether or not that vertex is the start vertex. Since all matrix elements between the computational basis state of that vertex and its neighbors are the same, one cannot distinguish the different neighbors in any way from the response to the given query. As we have mentioned, though, one can determine if a queried vertex is the start vertex since the Hamiltonian will have an additional coupling $t(0)\langle s|+h.c.)$.

Using the knowledge of which vertex is the start vertex it is not hard to distinguish the two graphs in O(1) queries: query the start vertex s to get some new vertex v, then query a neighbor of v (other than s, i.e. nonbacktracking) to get some new vertex w, then finally query a neighbor of w, again without backtracking. For C, with probability 1/3 that neighbor of w will be the start vertex. On the other hand, for D, the neighbor will never be the start vertex.

Suppose however, that we use only the information about the degree of the vertex and not which vertex is the start vertex. In this case, it is impossible to distinguish C from D using fewer than h queries because, trivially, any vertex accessed with fewer than h queries is not a leaf of D and hence has the same degree (i.e., 3) as every vertex in C, and hence the terms in the Hamiltonian coupling the corresponding computational basis state to its neighbors are the same.

So, if we choose h superpolynomially large, then item **4** is "almost satisfied", i.e., without information about the start vertex we cannot distinguish them with polynomially many queries. However, if we choose h superpolynomially large, then item **5** is not satisfied since the number of vertices is not $O(2^{\text{poly}(N)})$.

If instead we choose h only polynomially large, there is a simple efficient classical algorithm to distinguish C from D even without using knowledge of the start vertex: since we can avoid backtracking in the given query model, we will arrive at a leaf in h queries. Even if we perform a random walk on D, allowing backtracking, we will typically arrive at a leaf in O(h) queries

5 Decorated Graphs

In this section we define an operation called *decoration* and then define graphs C, D in terms of this operation. Our decoration operation is very similar to (indeed, it is a special case of) the decoration defined in [19] (there are other uses of the term "decoration" in the math literature, such as in set theory, which are unrelated to this). These graphs C, D will depend upon a large number of parameters; in Section 7 we will show that for appropriate choice of these parameters, all properties in Table 1 are fulfilled.

Section 5.1 defines decoration. The idea of decoration is to make it easy to for a classical algorithm to "get lost". We will added additional vertices and edges to some graph in decorating and a classical algorithm will tend to follow what one may call "false leads" along these edges so that it is hard for it to determine properties of the graph before decoration because it takes a large number of queries to avoid these false leads.

Section 5.2 applies decoration to define C, D and explains some motivation for this choice of C, D. Section 5.3 considers the spectrum of the adjacency matrices of C, D, as well as proving some properties of the eigenvector of the adjacency matrix of C with largest eigenvalue. Here we, roughly speaking, bound the effect of decoration on the spectrum and leading eigenvector of the graph.

5.1 Decoration

In this subsection, we define an operation that we call *decoration* that maps one graph to another graph.

We first recall some graph theory definitions. For us, all graphs are undirected, so all edges are unordered pairs of vertices, and there are no multi-edges or self-edges so that the edge set is a set of unordered pairs of distinct vertices. An m-ary tree is a rooted tree (i.e., one vertex is referred to as the root) in which each vertex has at most m children. A full m-ary tree is a tree in which every vertex has either or 0 or m children (i.e., all vertices which are not leafs have m children). A perfect m-ary tree is a full m-ary tree with all leaves at the same distance from the root. A binary tree is an m-ary tree with m=2.

We will make an additional definition (this concept may already be defined in the literature but we do not know a term for it). We will say that an inner regular graph is a graph such that there exists some vertex set T so that every vertex not in T has has degree d or has degree 1. We will say that such a graph is d-inner regular and we will say that T is the set of "terminal vertices" of the graph. The set of vertices not in T will be called the inner vertices.

We define the height of a tree to be the length of the longest path from the root to a leaf, so that tree with just one vertex (the root) has height 0 (sometimes it is defined this way in the literature, but other authors define it differing by one from our definition). Then, the number of vertices in a perfect m-ary tree of height h is $1 + m + m^2 + \ldots + m^h = (m^{h+1} - 1)/(m-1)$.

We now define decoration. If we (c, m, h)-decorate a graph G, the resulting graph is given by attaching c perfect m-ary trees of height h-1 to each vertex of G; that is, for each vertex v of G, we add c such trees, adding one edge from v to the root of each tree, so that the degree of v is increased by c. Call the resulting graph H. Then, any vertex of graph G which has degree d corresponds to a vertex of H with degree d+c. We may regard G as an induced subgraph of H; if G is a rooted tree, then the root of G corresponds to some vertex in H that we will regard as the root of H. We will refer to those vertices of H which correspond to some vertex of G as the original vertices of H, i.e., the original vertices of H are those in the subgraph G. If G is a full n-ary tree, and if we (n, 2n, h) decorate G, then H is a full (2n)-ary tree.

Now we define a sequence of decorations. Consider some sequence of heights h_1, h_2, \ldots, h_l for some l. Given a graph G_0 which is (n+1)-inner regular, we $(n, 2n, h_1)$ -decorate G, calling the result G_1 . The terminal set of G_1 will be the set of vertices which correspond to vertices in the terminal set of G, as well as additional leaves (vertices of degree 1) added in decoration, so that G_1 is (2n+1)-inner regular. We then $(2n, 4n, h_2)$ -decorate G_1 , calling the result G_2 , defining the terminal set of G_2 in the

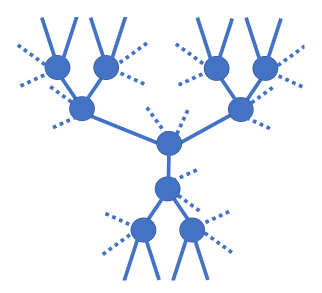


Figure 1: Example of decoration. Solid lines are edges of graph G which is 3-inner regular. In the notation given, it is $T_{3,2}$. Dashed lines represent edges added after (2,4,1)-decorating G. Solid circles are inner vertices and terminal vertices are not shown. Since $h_1=1$, the 4-ary trees added are of height 0; if we had taken $h_1=1$, then each of the dashed lines shown would have a solid circle and four additional dashed lines attached to them.

analogous way. Proceeding in this fashion, we $(2^m \cdot n, 2^{m+1} \cdot n, h_m)$ -decorate G_m , giving G_{m+1} , until we have defined graph G_l . We say that G_l is given by decorating G with height sequence h_1, \ldots, h_l . Note that each graph G_m is inner regular.

We will call the *original vertices* of G_l those vertices which correspond to some vertex of G_0 in the obvious way, i.e., G_0 is an induced subgraph of G_k and the original vertices are the vertices in that subgraph.

Remark: the reason that we talk about (n+1)-inner regular, rather than n-inner regular is that if one avoids backtracking, this means that there are n, rather than (n-1), choices of vertex to query from any given vertex. Also, since we are decorating with trees, in this way n refers to the arity of the tree rather than the degree of the tree. This is just an unimportant choice of how we define things and other readers might find it more convenient to shift our value of n by one.

For use later, let us define T_{n,h_0} where T_{n,h_0} is constructed by attaching n+1 perfect n-ary trees to some given vertex, i.e., T_{n,h_0} is n-inner regular with the terminal set being the leaves of the trees. See Fig. 1 for an example. Here we define

5.2 Application of Decoration, and Universal Cover of Graph

We will apply this decoration to two different choices of (n + 1)-inner regular graphs, each of which has some fixed vertex that we call the start vertex.

In the first case, we pick G_0 to have vertices labelled by a pair of integers (i, j) with $0 \le i < L$ for some L > 1 and $0 \le j < m$ for m = (n + 1)/2 for some odd n. There is an edge between (i, j) and (k, l) if $i = k \pm 1 \mod L$; note that there is no constraint on j, l. So, for m = 1, the graph is a so-called ring graph. The start vertex will have $i = \lfloor L/2 \rfloor$ and j = 0. We denote this graph $R_{L,m}$ where m = (n + 1)/2.

The second case is the same except that there is an edge between (i,j) and (k,l) if $i=k\pm 1$. Note that the " mod L" is missing in the definition of D. Again the start vertex has $i=\lfloor L/2\rfloor$ and j=0. So, for m=1, this graph is a so-called path graph or linear graph. We denote this graph $P_{L,m}$.

See Fig. 2.

Define $C = G_l$ in the case that $G = R_{L,(n+1)/2}$ and define $D = G_l$ in the case that $G = P_{L,(n+1)/2}$.

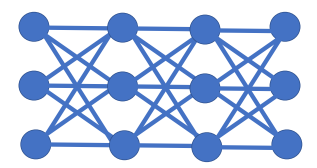


Figure 2: Example of graph $P_{4,3}$. Solid circles are vertices and solid lines are edges. Note that the graph is 6-inner regular so n=5. In the case of graph $R_{4,3}$, there are 9 additional edges connecting each of the three vertices at the left to each of the three vertices on the right.

Note that $R_{L,(n+1)/2}$ is an (n+1)-regular graph and $P_{L,(n+1)/2}$ is an (n+1)-inner regular graph where the terminal vertices are those (i,j) with i=0 or i=L-1.

Then, in both cases, the graph G_l is a d-inner regular graph with $d = 2^{l+1} \cdot n$. We will define a start vertex on G_l in the obvious way: it is the original vertex in G_l that corresponds to the start vertex of G.

The key then is that to distinguish C and D, one must be able to go a long distance, of order L, on the graph. Decoration will make it hard for any classical algorithm to follow such a long path.

5.3 Spectrum of Adjacency Matrix of Decorated Graph

We now provide bounds on the spectrum of the adjacency matrix of graph G_l constructed from decorating G with some height sequence. For any graph H, let $\lambda_0(H)$ denote the largest eigenvalue of the adjacency matrix of H. Since G is a subgraph of G_l , it follows that $\lambda_0(G_l) \geq \lambda_0(G)$.

We have:

Lemma 4. Assume $n/L^2 = \omega(1)$.

For either choice of G_0 , the largest eigenvalue of the adjacency matrix of G_l is bounded by

$$\lambda_0(G) \le \lambda_0(G_l) \le \lambda_0(G) + 2^{l/2+1} \cdot \sqrt{n}$$
.

Further $\lambda_0(C) \geq n$ and $\lambda_0(D) \leq n + 2^{l/2+1} \cdot \sqrt{n} - \Theta(n/L^2)$.

The second largest eigenvalue of the adjacency matrix of C is upper bounded by $n + 2^{l/2+1} \cdot \sqrt{n} - \Theta(n/L^2)$.

Finally, let ψ be an eigenvector of the adjacency matrix of C with largest eigenvalue (the eigenspace has dimension 1 since G_l is connected), with $|\psi| = 1$. Let Π be a diagonal matrix which is 1 on the original vertices of C and 0 on the other vertices. Let $p = |\Pi\psi|^2$. Remark: heuristically, p is the "probability" that if one measures ψ in a basis of the vertices, that the result will be one of the original vertices. Then, if $n \geq 2^{l/2+2} \cdot \sqrt{n}$, $p \geq 1/5$.

Proof. To show the first result, let A be the adjacency matrix of G_l . We decompose $A = A_0 + A_1$ where $A_0 = \Pi A \Pi$ so that A_0 is the adjacency matrix of the subgraph of G_l obtained by deleting all edges except those which connect two original vertices. Then, $||A_0|| = \lambda_0(G)$ and so $\lambda_0(G_l) \leq \lambda_0(G) + ||A_1||$.

However, A_1 is equal to the adjacency matrix of the subgraph of G_l obtained by deleting any edge connecting two original vertices of G_l . This subgraph is a forest, to use the terminology of graph theory: it consists of disconnected trees, one tree for each vertex in G. Indeed each of these trees is an m-ary tree with $m = 2^l \cdot n$ (recall that if G has degree n + 1, then G_l has degree $2^l \cdot n + 1$).

We now upper bound the largest eigenvalue of the adjacency matrix of an m-ary tree. For a perfect m-ary tree of height h, it is possible to compute the largest eigenvalue: the tree has a symmetry

under permuting the daughters of any given vertex and the largest eigenvector will be invariant under this symmetry. So, let v_k , for integer k with $0 \le k \le h$, denote the vector with norm 1 which is an equal amplitude superposition of all vertices which are distance k from the root, so that v_0 is 1 on the root and 0 elsewhere, v_1 has amplitude $1/\sqrt{m}$ on each of the daughters of the root, and so on. We can then write the adjacency matrix, restricted to the subspace spanned by v_0, v_1, \ldots as $\sqrt{m} \sum_{0 \le k \le k+1} |v_k\rangle \langle v_{k+1}|| + h.c.$ and so clearly the largest eigenvalue is bounded by $2\sqrt{m}$. So, $\lambda_0(G_l) \le \lambda_0(G) + 2^{l/2+1} \cdot \sqrt{n}$, as claimed

The lower bound on $\lambda_0(C) \ge n$ follows from the fact that $\lambda_0(R_{L,(n+1)/2}) = n+1 > n$. The upper bound on $\lambda_0(D)$ follows since $\lambda_0(P_{L,(n+1)/2}) = n+1 - \Theta(n/L^2) = n - \Theta(n/L^2)$ since $n/L^2 = \omega(1)$.

To bound the second largest eigenvalue of the adjacency matrix of C, again decompose $A = A_0 + A_1$. By the Courant-Fischer-Weyl min-max principle, the second largest eigenvalue of A is equal to the minimum, over all subspaces of codimension 1, of the largest eigenvalue of the projection of A into that subspace. Consider the subspace orthogonal to the eigenvector of A_0 of largest eigenvalue; the projection of A_0 into this subspace is bounded by $n - \Theta(n/L^2)$ and so the second largest eigenvalue of A is upper bounded by $n - \Theta(n/L^2) + ||A_1||$.

For the final claim, we have $\lambda_0(C) \leq p||A_0|| + 2\sqrt{p(1-p)}||A_1|| + (1-p)||A_1||$. Since $\lambda_0(C) \geq n$ and $||A_0|| = n$, we have $(1-p)n \leq \left((1-p) + 2\sqrt{p(1-p)}\right)2^{l/2+1} \cdot \sqrt{n+1}$. So, for $n \geq 2(2^{l/2+1} \cdot \sqrt{n})$, after a little algebra we find that $p \geq 1/5$.

6 Classical Hardness

We now show classical hardness. We will give a lower bound on the number of queries needed by a classical algorithm to distinguish C from D, with the initial state of the classical algorithm being the start vertex. The lower bound Lemma 5 will depend on the difficulty of reaching a certain set Δ defined below. To show this difficulty, the main result is in the inductive Lemma 6; we then apply this lemma in Section 7.

Given a d-inner regular graph, with some given choice of start vertex, and given some set S which is a subset of the set of vertices, we say that it is (p,q)-hard to reach S if no classical algorithm, starting from the start vertex, can reach S with at most q queries with probability greater than p. Here "reach" means that at some point the classical algorithm queries a vertex in S.

Note that given a perfect m-ary tree of height h, it is clearly (0, h + 1)-hard to reach the set of leaves of the tree if the root vertex is the start vertex: the first query will query the start vertex, the second query can query a vertex at most distance 1 from the start vertex, and so on.

Our goal will be to show for graph D that it is hard (for some choice of parameters p, q) to reach the set Δ of vertices of D which correspond to terminal vertices of G_0 , i.e., Δ is a subset of the terminal vertices of G_l , consisting only of those which correspond to terminal vertices of G_0 rather than leaves added in decoration.

This will then imply a bound on the ability of a classical algorithm to distinguish C from D in the modified query model:

Lemma 5. Suppose it is (p,q)-hard to reach the set Δ above for graph D. Suppose one chooses some graph G to be C or D with probability 1/2 for each choice. Then, no classical algorithm using at most q queries in the modified query model can correctly guess which graph G is with probability greater than 1/2 + p/2.

Additionally, the mutual information between the random variable G and the query responses is $\leq p$.

Proof. Consider the set of vertices in C which correspond to an original vertex (i, j) with i = 0 or i = L - 1. In an overload of notation, let us also call this set Δ .

Consider some classical algorithm. If we apply this algorithm to graph D or graph C, in the modified query model the two algorithms have the same probability distribution of query responses conditioned on the case that the algorithm applied to D does not reach Δ .

Hence, it is (p,q)-hard to reach the set Δ in C.

So, for graph D, the probability distribution of query responses is $(1-p)\sigma+p\tau$ for some probability distribution τ : the first term in the sum is the case conditioned on not reaching Δ and the second is the case when it reaches Δ . For C, the probability distribution of query responses is $(1-p)\sigma+p\mu$ for some probability distribution μ .

The ℓ_1 distance between the two probability distributions is at most 2p. So, the probability that the algorithm guesses right is at most 1/2 + p/2 (to see this, let P, P' be two probability distributions on some set of events; let e label events; choose P or P' with probability 1/2, then observe some event given the probability distribution; let $A_e = 1$ if one choose P on event e and $A_e = 0$ otherwise; then the probability of guessing right is $\sum_e A_e P_e/2 - \sum_e (1 - A_e) P'_e/2 = 1/2 + \sum_e (A_e - 1/2) (P_e - P'_e)/2$ and $\sum_e (A_e - 1/2) (P_e - P'_e)/2 \le |P - P'|_1/4$).

The claim about the mutual information follows because the optimal case for the mutual information is when the support of σ , τ , μ are all disjoint.

We now need a couple more definitions. First,

Definition 1. We will assign a number called the level to each vertex v of any graph G_k defined by some sequence of decorations of a graph as follows. Decoration gives a sequence of graphs G_0, G_1, \ldots, G_k . For j < k, we may regard G_j as an induced subgraph of G_k . The level of a vertex $v \in G_k$ is equal to the smallest j such that $v \in G_j$. Hence, the original vertices of G_k are those with level 0.

Second, consider the problem of tossing a coin which is heads with probability 1/2 and tails with probability 1/2. Define $P_{coin}(n, N)$ to be the probability that one observes n heads after at most N tosses of the coin. This is the same as the probability that, after tossing the coin N times, one has observed at least n heads. Thus,

$$P_{coin}(n,N) = \sum_{m=n}^{N} 2^{-N} \binom{N}{m}.$$

Now we give a lemma with two parts. This first part is an inductive lemma that implies difficult of reaching leaves of a graph obtained by decorating a tree $T_{n,h}$. The second part is used to prove difficult of reaching Δ ; both parts have very similarly proofs and in applications we will first apply the first part inductive for several height sequences and then use the result as an input to the second part of the lemma. Throughout, we will use S_{leaf} to refer to a set of leafs in a tree graph; this will be the set of terminal vertices.

Now we show:

Lemma 6. Assume that for some n, h_0, h_1, \ldots, h_k , it is (P,Q)-hard to reach S_{leaf} in graph G given by decorating T_{2n,h_0} by height sequence h_1, \ldots, h_k . Then:

1. For any integers $M, H \geq 0$, it is (P', Q')-hard to reach S_{leaf} in graph G' given by decorating $G = T_{n,H}$ with height sequence h_0, h_1, \ldots, h_k , where

$$P' = P_{coin}(H - h_0, M) + Q'P, (4)$$

and

$$Q' = (M - (H - h_0))Q. (5)$$

2. For any integers $M, L \ge 0$, it is (P', Q')-hard to reach Δ in graph D given by decorating $P_{L,(n+1)/2}$ with height sequence h_0, h_1, \ldots, h_k where

$$P' = P_{coin}(L/4 - h_0 - O(1), M) + Q'P,$$
(6)

and

$$Q' = (M - (H - h_0))Q. (7)$$

Proof. We prove the first claim first. After q queries by the algorithm, we can describe the queries by a tree T(q) with q edges, each vertex of which corresponds to some vertex in G'. We use letters a, b, \ldots to denote vertices in this tree T(q) and use v, w, \ldots to vertices in G'. The root of the tree T(q) corresponds to the start vertex. If some vertex $a \in T(q)$ corresponds to some $v \in G'$, then the daughters of a correspond to neighbors of v obtained by querying v.

Given an $a \in T(q)$, we will say it has some given level if the corresponding vertex in G' has that level. For any vertex $a \in T(q)$ or any $v \in G'$, we say that the *subtree* of a (respectively, v) is the tree consisting of a (respectively, v) and all its descendants. "Queries of a subtree" mean queries in the modified query model starting from the root of that subtree.

Define a subtree S(q) of T(q). S(q) will be the induced subgraph whose vertices consist of the root of T(q) and of all other vertices a of T(q) which have been queried at least Q times and such that the parent of the given vertex a is at level 0.

Suppose after some number of queries q, some new vertex a is added to S(q). We will now consider the probability distribution of the level of a, conditioned on the level being 0 or 1. Let a correspond to vertex v of G'. Suppose that v has distance at most $H - h_0$ from the root of G' so that S_{leaf} in G' is distance at least h_0 from v.

We will say that a vertex in G' has property (\dagger) if these three conditions hold: it is distance $\leq H - h_0$ from the root of G', and is level 0 or 1, and is the the child of a vertex with level 0. For a v with property (\dagger) , we will say that event (*) occurs for that v if we reach a vertex in the subtree of v which is level 0 or 1 and distance $\geq h_0$ from v with fewer than Q queries in that subtree. If v has level 1, then the responses to queries of the subtree of v are the same as in the given query model on graph G given by decorating T_{2n',h_0} by height sequence h_1,\ldots,h_k . In that case, event (*) occurs iff we reach S_{leaf} in G in fewer than Q queries; this probability is bounded by P by the inductive assumption. If instead v has level 0, then since by assumption S_{leaf} in G' is distance at least h_0 from v, the probability of event (*) occurring for that v is also bounded by P. The key point here is that if v has level 0 and we consider the subtree of v, and then further consider the subgraph of that subtree consisting of vertices of distance $\leq h_0$ from v, this subgraph is isomorphic to G.

Now, let us condition on event (*) not occurring for any v. At the end of the proof of the lemma, we will upper bound the probability of event (*) occurring. If event (*) does not occur, then the distribution of queries responses in the subtree of v is the same whether v has level 0 or 1. So, when vertex a is added to S(q), it has probability 1/2 of being 0, conditioned on the level being 0 or 1, i.e., we toss an unbiased coin to determine the level of that vertex: "heads" corresponds to level 0 and "tails" corresponds to level 1. The level may also be > 1, but including that possibility only increases the number of queries.

The number of queries q is at least equal to Q times the number of "tails" that have occurred. Remark: "heads" also implies that there were at least Q queries of the subtree of some vertex, but if we include those queries due to "heads" in the total number, we must be careful to avoid overcounting as those Q queries of the subtree of some vertex v_i will also give some number of queries (up to Q-1) of descendants of v_i . So, for simplicity, we will use the lower bound that the number of queries is at least Q times the number of tails.

Starting from the root of G', to reach S_{leaf} in G' with at most q queries, we must have one of these two possibilities: (1) subtree S(q) contains some vertex at level 0 with distance $\geq H - h_0$ from the root; or (2) we reach S_{leaf} in G' in fewer than Q queries starting with some vertex v of level 0 and distance $\leq H - h_0$ from the root, i.e., we reach S_{leaf} in G' in the subtree of such a vertex v with fewer than Q queries in that subtree.

Conditioned on event (*) not occurring, the probability of 1 occurring in at most $(M - (H - h_0))Q$ queries s bounded by the probability of having at least $H - h_0$ heads out of M coin tosses and so is bounded by $P_{coin}(H - h_0, M)$. If event 2 happens, then event (*) happens. By a union bound, since there are only Q' vertices that we query, event (*) happens with probability at most Q'P. So, by a union bound, the probability of reaching S_{leaf} in G' with at most QM queries is bounded by $P_{coin}(H - h_0, M) + Q'P$.

Remark: likely the union bound in the previous paragraph on the probability of event (*) could

be tightened. If we query a vertex at distance $H - h_0$ from the root then indeed the probability of reaching S_{leaf} is bounded by P but for vertices of lower distance from the root the probability is less as would follow from a better inductive assumption. We will not need this tightening so we omit it.

Having proved the first claim, the proof of the second claim is almost identical. We use two new ideas. The set Δ is at distance L/2 - O(1) from the start vertex of D. So, to reach Δ one must at some time query some vertex v which is at level 0 and at distance $\lfloor L/4 \rfloor$ from the start vertex of D such that in the subtree of v one then queries a vertex in Δ .

This introduction of vertex v is the first new idea: by choosing such a vertex v which is far from the start vertex, we will be able to ignore the possibility that the algorithm gets extra information about which vertex is the start vertex in response to queries by considering only vertices near v. Indeed, consider queries in the subtree of v. Consider the set of vertices of D including all vertices of level ≥ 1 and all vertices of level 0 which are distance L/4-O(1) so that this set does not intersect Δ and does not contain the start vertex. Let D' be the subgraph of D induced by this set of vertices. In the modified query model one cannot distinguish D' from its universal cover \tilde{D}' , which is a tree graph; this is the second new idea. Take the root of this tree graph to have its image under the covering map be v. Then, responses to queries on this tree graph are the same as almost the same in queries on the graph given by decorating $T_{n,L/4-O(1)}$ h_0, h_1, \ldots, h_k . The only minor difference is that we may take the first query to be nonbacktracking, so that rather than decorating graph $T_{n,L/4-O(1)}$ we decorate an n-ary tree of depth L/4-O(1), i.e., there are n rather than n+1 new neighbors in response to the first query. To reach Δ , one must reach some vertex which is at level 0 and at distance L/4-O(1) from v.

Then, the rest of the proof is the same.

7 Choice of Height Sequence and Proof of Main Theorem

We now make specific choices of the height sequence to prove Theorem 1.

We pick $n = N^8$ in the construction of C, D. The value of n does not matter for the classical lower bounds that follow from Lemma 6. However, the choice of n does affect the spectrum of the quantum Hamiltonian. We pick L = 4(l+1)N + O(1). Thus $n/L^2 = \Theta(N^6/l^2)$.

We decorate with height sequence h_1, \ldots, h_l , choosing $l = \lfloor \log_2(N) \rfloor$. Thus, the graph D is d-inner regular, with $d = \Theta(N^9)$. We pick

$$h_k = N \cdot (l+1-k),$$

for that $h_{k-1} - h_k = N$ and $L/4 - O(1) - h_1 = N$.

We first quickly show items **1-3,5** of Table 1 before showing the harder result, item **4**. Then, Theorem 1 follows from Lemma 3.

Note that $n/L^2 \gg \sqrt{d}$. So, from Lemma 4, it follows that the difference between the ground state energy of C and D is $\Theta(N/L^2)$ and so item 1 of Table 1 is satisfied.

Again since $n/L^2 \gg \sqrt{d}$, from the bound on the second largest eigenvalue of the adjacency matrix of C in Lemma 4, we satisfy the condition on the spectral gap of the Hamiltonian of C, item 2 of Table 1.

Item 3 of Table 1 is also satisfied by the last result in Lemma 4, since there are only O(N) original vertices and the amplitude of the ground state wavefunction is the same on all of them.

Item 5 of Table 1 trivially follows. The number of vertices in G_0 is $\leq \text{poly}(N)$. Decoration by an m-ary tree of height h multiplies the number of vertices in the graph by $O(m^h)$. All h_k are $O(N \log(N))$, and the largest m is O(poly(N)), and there are only $O(\log(N))$ steps of decoration, so the total number of vertices is $\exp(O(N \log(N)^3))$.

Remark: since the construction of an admissible path in Section 3 uses N copies of the graph, the number of computational basis states needed is $\exp(O(N^2 \log(N^3)))$. This number can be reduced somewhat since we could choose smaller values of h_k , such as $h_k = N^{\alpha} \cdot (l+1-k)$ for some smaller $\alpha < 1$; we omit this.

To prove item 4, we use Lemma 6. We first use the first part of the lemma. Consider a sequence of graphs $J_l, J_{l-1}, J_{l-2}, \ldots, J_1$, where J_k is given by decorating T_{n_k, h_k} by height sequence $h_{k+1}, h_{k+2}, \ldots, h_l$, for $n_k = n \cdot 2^k$.

Clearly, for J_l , it is (0, N-1)-hard to reach S_{leaf} , simply because the tree has height N. We apply Lemma 6 to use (P_{k+1}, Q_{k+1}) -hardness on J_{k+1} to show (P_k, Q_k) -hardness on J_k . For all k, we pick M = (4/3)N, so that

$$Q_k = (N/3)^{l-k}N. (8)$$

Then $P_{coin} = \exp(-\Omega(N))$ and Eq. (4) gives that $P_{k-1} \leq \exp(-\Omega(N)) + NP_k$. Hence

$$P_{k} \leq \exp(-\Omega(N)) + Q_{k}P_{k+1}$$

$$\leq \exp(-\Omega(N)) + Q_{k}\exp(-\Omega(N)) + Q_{k}Q_{k+1}P_{k+2}$$

$$\leq \dots$$

$$\leq \exp(-\Omega(N))(1 + Q_{k} + Q_{k}Q_{k+1} + \dots)$$

$$\leq \exp(-\Omega(N))O(Q_{k}^{l})$$

$$\leq \exp(-\Omega(N))\exp(O(\log(N)^{3}))$$

$$\leq \exp(-\Omega(N)).$$

$$(9)$$

It is (P,Q)-hard to reach S_{leaf} in J_1 with

$$P = \exp(-\Omega(N)),\tag{10}$$

and

$$Q = \exp(\Theta(\log(N)^2)). \tag{11}$$

Then, use the second part of the lemma, with the same M, to show

Lemma 7. It is (P,Q)-hard to reach Δ in D with

$$P = \exp(-\Omega(N)),\tag{12}$$

and

$$Q = \exp(\Theta(\log(N)^2)). \tag{13}$$

This completes the proof of Theorem 1.

We make one final remark on the Hamiltonian on these decorated graphs. For the given parameters, the ground state wavefunction on C has most of its probability (its ℓ_2 norm) on the original vertices of C. However, the ℓ_1 norm is concentrated near the terminal vertices. The distinction between ℓ_1 and ℓ_2 norm was used in [14] to "pin" the worldline in the case of path integral Monte Carlo with open boundary conditions and was considered in [20] as an obstruction for diffusion Monte Carlo methods. The large ℓ_2 norm of the ground state wavefunction on C can be regarded as arising from all the short cycles on C; if we replace C with a (finite) tree with the same degree, then ground state energy on C shifts and the ℓ_2 norm of the ground state becomes concentrated instead near the terminal vertices. Thus, one may say that the topological obstructions are related to the ℓ_1 versus ℓ_2 obstructions. The idea of our construction here is to make it so that no classical algorithm can (once one is far from the start vertex) efficiently distinguish original vertices from other vertices, since it cannot detect the short cycles between the original vertices and so it is unable to determine that the ℓ_2 norm should be larger.

8 Linear Paths

We have considered a path H_s given by an oracle and we have implemented some complicated s-dependent terms so that certain terms increase and then later decrease over the path. One might be

interested in the case of linear interpolation so that $H_s = (1 - s)H_0 + sH_1$ for some fixed and known H_0 , such as a transverse field, with H_1 being a diagonal matrix given by an oracle. It seems likely that the construction here could be adapted to show hardness in that case too, using some perturbative gadgets. Since it is not too interesting a question, we only sketch how this might be done.

One might induce dynamics on an unknown graph (either C or D) as follows. Both graphs have the same vertex set but have different edge sets. Define a new graph E which has one vertex for each vertex in C and one vertex for each edge in C. Given a pair of vertices v, w connected by an edge e in C, the graph E will have an edge between the vertex corresponding to v and the vertex corresponding to e, as well as an edge between the vertex corresponding to w and the vertex corresponding to e. We add also a diagonal term to the Hamiltonian on graph E which assigns some positive energy to all vertices which correspond to edges of C. If all these diagonal terms are the same and are chosen appropriately, we have a perturbative gadget which gives us an effective Hamiltonian corresponding to the Hamiltonian of graph C, up to an overall multiplicative scalar. On the other hand, we could increase the diagonal term on vertices which correspond to edges of C which are not edges of D to effectively induce the Hamiltonian on D.

Also, in our construction of a path in Section 3 we used the ability to turn on and off a term $t|0\rangle\langle s|+h.c.$ in the Hamiltonian. Such a term it seems can be induced by some perturbative gadgets also. First consider a slightly more general case where $H_0 = (1-s)H_0 + H_s^{diag}$ where H_0 is fixed, known term and H_s^{diag} is a diagonal matrix depending on s and given by an oracle. Then, we can effectively induce a t which depends on s by using hopping between an intermediate state, i.e., $-(|0\rangle\langle \text{int}| + |\text{int}\rangle\langle s|) + h.c.$ where $|\text{int}\rangle$ is some intermediate state. Then adding an s-dependent diagonal term on $|\text{int}\rangle$ can be used to turn on or off the effective hopping between $|0\rangle$ and $|s\rangle$.

To effectively induce this s-dependent diagonal term on $|\text{int}\rangle$ we could use another trick: replace every basis state with some set of basis states of size poly(N) and adding hopping terms between every pair of basis states in each such set. Then, each such set defines a subspace. Further add some diagonal terms on some of the basis states in each set, so that, for example, one set might include n_0 states with energy E_0 , some number $n_1 \gg n_0$ of additional states with energy $E_1 \gg E_0$, and so on. Then, if s is close to 1 so that $(1-s)H_0$ is small, almost of the amplitude will be in the states with energy 0 and we can treat that set of states as a single state with energy $E_0 - (1-s)(n_0-1)$; for slightly smaller s, we will start to occupy higher energy states and for some choices of sequences n, E we can treat that set approximately as a single state with energy $\approx E_1 - (1-s)(n_0+n_1-1)$, and so on. By adjusting the sequences of n, E on each set, it seems that we can effectively implement a problem with fairly complicated s-dependent diagonal terms.

References

- [1] Edward Farhi, Jeffrey Goldstone, Sam Gutmann, Joshua Lapan, Andrew Lundgren, and Daniel Preda. A quantum adiabatic evolution algorithm applied to random instances of an np-complete problem. *Science*, 292(5516):472–475, 2001. doi:10.1126/science.1057726.
- [2] Boris Altshuler, Hari Krovi, and Jeremie Roland. Adiabatic quantum optimization fails for random instances of np-complete problems. arXiv preprint arXiv:0908.2782, 2009.
- [3] Boris Altshuler, Hari Krovi, and Jérémie Roland. Anderson localization makes adiabatic quantum optimization fail. *Proceedings of the National Academy of Sciences*, 107(28):12446–12450, 2010. doi:10.1073/pnas.1002116107.
- [4] Sergey Knysh and Vadim Smelyanskiy. On the relevance of avoided crossings away from quantum critical point to the complexity of quantum adiabatic algorithm. arXiv preprint arXiv:1005.3011, 2010.
- [5] M. B. Hastings. A short path quantum algorithm for exact optimization. *Quantum*, 2:78, jul 2018. URL: https://doi.org/10.22331%2Fq-2018-07-26-78, doi:10.22331/q-2018-07-26-78.

- [6] M. B. Hastings. The short path algorithm applied to a toy model. Quantum, 3:145, may 2019. URL: https://doi.org/10.22331%2Fq-2019-05-20-145, doi:10.22331/q-2019-05-20-145.
- [7] CR Laumann, R Moessner, A Scardicchio, and Shivaji Lal Sondhi. Quantum adiabatic algorithm and scaling of gaps at first-order quantum phase transitions. *Physical review letters*, 109(3):030502, 2012. doi:10.1103/physrevlett.109.030502.
- [8] Dave Wecker, Matthew B. Hastings, and Matthias Troyer. Training a quantum optimizer. *Physical Review A*, 94(2), aug 2016. URL: https://doi.org/10.1103%2Fphysreva.94.022309, doi: 10.1103/physreva.94.022309.
- [9] Troels F Rønnow, Zhihui Wang, Joshua Job, Sergio Boixo, Sergei V Isakov, David Wecker, John M Martinis, Daniel A Lidar, and Matthias Troyer. Defining and detecting quantum speedup. Science, 345(6195):420-424, 2014. doi:10.1126/science.1252319.
- [10] D. Aharonov, W. van Dam, J. Kempe, Z. Landau, S. Lloyd, and O. Regev. Adiabatic quantum computation is equivalent to standard quantum computation. In 45th Annual IEEE Symposium on Foundations of Computer Science. IEEE. doi:10.1109/focs.2004.8.
- [11] Andrew M. Childs, Edward Farhi, and Sam Gutmann. *Quantum Information Processing*, 1(1/2):35–43, 2002. doi:10.1023/a:1019609420309.
- [12] Rolando D. Somma, Daniel Nagaj, and Mária Kieferová. Quantum speedup by quantum annealing. *Physical Review Letters*, 109(5), jul 2012. doi:10.1103/physrevlett.109.050501.
- [13] Andrew M. Childs. Universal computation by quantum walk. *Physical Review Letters*, 102(18), may 2009. doi:10.1103/physrevlett.102.180501.
- [14] M. B. Hastings and M. H. Freedman. Obstructions to classically simulating the adiabatic algorithm. *QIC*, 13:1038, 2013.
- [15] Patrik Henelius, S. M. Girvin, and Anders W. Sandvik. Role of winding numbers in quantum monte carlo simulations. *Physical Review B*, 57(21):13382–13385, jun 1998. doi:10.1103/physrevb.57.13382.
- [16] Dorit Aharonov and Amnon Ta-Shma. Adiabatic quantum state generation and statistical zero knowledge. In *Proceedings of the thirty-fifth ACM symposium on Theory of computing 03*. ACM Press, 2003. doi:10.1145/780542.780546.
- [17] Andrew Macgregor Childs. Quantum information processing in continuous time. PhD thesis, Massachusetts Institute of Technology, 2004.
- [18] Dominic W. Berry, Graeme Ahokas, Richard Cleve, and Barry C. Sanders. Efficient quantum algorithms for simulating sparse hamiltonians. *Communications in Mathematical Physics*, 270(2):359–371, dec 2006. doi:10.1007/s00220-006-0150-x.
- [19] Jeffrey H. Schenker and Michael Aizenman. Letters in Mathematical Physics, 53(3):253–262, 2000. doi:10.1023/a:1011032212489.
- [20] Michael Jarret, Stephen P. Jordan, and Brad Lackey. Adiabatic optimization versus diffusion monte carlo methods. *Physical Review A*, 94(4), oct 2016. doi:10.1103/physreva.94.042318.