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Quantum-Merlin-Arthur–complete problems for stoquastic Hamiltonians and Markov matrices

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We show that finding the lowest eigenvalue of a 3-local symmetric stochastic matrix is Quantum-Merlin-Arthur-complete (QMA-complete). We also show that finding the highest energy of a stoquastic Hamiltonian is QMA-complete and that adiabatic quantum computation using certain excited states of a stoquastic Hamiltonian is universal. Our results give a QMA-complete problem arising in the classical setting of Markov chains and adiabatically universal Hamiltonians that arise in many physical systems.

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I. INTRODUCTION

Quantum complexity theory is the study of the capabilities and limitations of computational devices operating according to the principles of quantum mechanics [1]. Because many of the classical constructs of computer science (e.g., circuits and clauses) are replaced by matrices, quantum complexity theory is sometimes referred to as matrix-valued complexity theory [2,3]. In addition to its intrinsic interest, this subject has many connections to issues of practical relevance to physical science, such as the difficulty with computing properties of quantum systems using either quantum or classical devices [4–6].

Perhaps the most basic classical complexity classes are P, the class of problems solved by a deterministic Turing machine in polynomial time, and NP (nondeterministic polynomial), the class of problems whose verification lies in P. It is widely believed, but not proven, that NP is strictly larger than P [7].

Because quantum mechanics only predicts probabilities of events, the classical deterministic classes are not the most natural place to start if one seeks their quantum generalizations. The probabilistic generalization of P is bounded-error probabilistic polynomial time (BPP), those problems solvable by a probabilistic Turing machine in polynomial time with bounded error [8]. The quantum generalization of this class is bounded-error quantum polynomial time (BQP), the class of problems solvable in polynomial time with bounded error on a quantum computer [1].

The classical probabilistic generalization of NP is the class MA [9]. This generalizes NP to problems whose verification is in BPP. MA stands for Merlin-Arthur. Merlin, who is computationally unbounded but untrustworthy, provides a proof that Arthur can verify using his BPP machine. The class MA possesses a quantum generalization to quantum-Merlin-Arthur (QMA) [10–12]. QMA may be intuitively understood as the class of decision problems that can be efficiently verified by a quantum computer.

Given a classical description of a decision problem x of length n, the prover, Merlin, provides a witness state |ψ⟩ to the verifier, Arthur. Arthur then performs a poly(n)-time quantum computation on the witness |ψ⟩ and either accepts or rejects. A problem is contained in QMA if, for all YES instances, there exists a witness causing Arthur to accept with probability greater than 2/3 and, for NO instances, there does not exist any witness that causes Arthur to accept with probability greater than 1/3. A problem X is said to be QMA-complete if it is contained in QMA and every problem in QMA can be converted to an instance of X in classical polynomial time.

Let us consider the following question: What is the ground-state energy of a quantum system? This question lies at the core of many areas of physical science, including electronic-structure theory and condensed-matter physics. In quantum complexity theory, this has been formalized (originally by Kitaev [11]; see also, for example, [13]) as the k-local Hamiltonian problem. For some systems, complexity-theoretic arguments suggest that efficient computation of the ground-state energy is likely to remain beyond reach [11,14].

A Hamiltonian H, acting on n qubits, is said to be k-local if it is of the form

\[ H = \sum_{s} H_s, \]

where each H_s acts on at most k qubits. Thus, for example, 1-local Hamiltonians consist only of external fields acting on individual qubits, and 2-local Hamiltonians consist of 1-local terms and pairwise couplings between qubits. Physically realistic Hamiltonians are usually k-local with small k, often 2 or 1, and each local term has bounded norm. Note that this notion of locality has nothing to do with spatial locality; a 2-local Hamiltonian may have long-range couplings, but they must be pairwise.
**Problem: k-local Hamiltonian**

**Input.** We are given a classical description of a $k$-local Hamiltonian $H$ on $n$ qubits $H = \sum_{j=1}^{r} H_j$ with $r = \text{poly}(n)$. Each $H_j$ acts on at most $k$ qubits and has $O(1)$ operator norm. In addition, we are given two constants $a$ and $b$ such that $0 \leq a < b$, and $b - a = \epsilon > 1/\text{poly}(n)$.

**Output.** If $H$ has an eigenvalue $\leq a$, answer YES. If all eigenvalues of $H$ are $> b$, answer NO.

**Promise.** The Hamiltonian is such that it will produce either YES or NO.

Perhaps a more obvious formulation of this problem is to ask for an approximate ground-state energy to within $\pm \epsilon$ of the correct answer. However, if one can decide the answer to $k$-local Hamiltonian in polynomial time, then one can solve the approximation version in polynomial time with a binary search. Thus, the approximation problem is of equivalent difficulty to the “decision” version to within a polynomial factor.

The problem $k$-local Hamiltonian is QMA-complete for $k \geq 2$ [13]. The $k$-local Hamiltonian problem is specified by the matrix elements of the local terms of $H$. YES instances possess the ground state as a witness. The verification circuit is the phase estimation algorithm—a suitably formalized version of the notion of energy measurement [11]. If the lowest eigenvalue of $H$ is less than $a$ (a YES instance), then Arthur will accept the ground state as a witness. However, if the lowest eigenvalue of $H$ is greater than $b$ (a NO instance), then Merlin cannot supply any eigenstate or superposition of eigenstates that will result in a measurement of energy less than $b$.

It is considered unlikely that QMA $\subseteq$ BQP, and therefore it is probably impossible to construct a general quantum (or classical) algorithm that finds ground-state energies in polynomial time. Thus, unless QMA $\subseteq$ AM (which is believed to be unlikely), such quantum computation is restricted to stochastic Hamiltonians. These are Hamiltonians in which all matrix elements are real and non-negative, and the sum of matrix elements in any row or column is 1.

**Problem: Quantum k-SAT**

**Input.** A set of $k$-local projectors $\{\Pi_q\}$ for $q \in \{1, \ldots, m\}$, where $m = \text{poly}(n)$ and a parameter $\bar{\epsilon} > 1/\text{poly}(n)$.

**Output.** If there is a state $|\phi\rangle$ such that $\Pi_1|\phi\rangle = 0$ for each $q \in \{1, \ldots, M\}$, then this is a YES instance. If every state $|\phi\rangle$ satisfies

$$\sum_{q=1}^{M} \langle \phi | \Pi_q | \phi \rangle \geq \bar{\epsilon},$$

then it is a NO instance.

**Promise.** The instance is either YES or NO.

In [2] the stoquastic restriction of quantum $k$-SAT was shown to be contained in MA for any constant $k$, and MA-complete for $k = 6$—the first nontrivial example of an MA-complete problem. In [3] these results were extended to a simplified form of stoquastic quantum $k$-SAT in which projectors $\Pi_q$ all have matrix elements taken from the set $\{0, 1/2, 1\}$, and the stoquastic constraints which appear as terms in the Hamiltonian are of the form $H_q = I - \Pi_q$.

The main intuition behind these results is that, by the Perron-Frobenius theorem, the ground state of a stoquastic Hamiltonian consists entirely of real positive amplitudes (given the appropriate choice of global phase). Thus, the ground state is proportional to a classical probability distribution. For this reason, ground-state properties are amenable to classical random-walk algorithms and certain problems such as stoquastic $k$-local Hamiltonian fall into classical probabilistic complexity classes such as AM. Diffusion quantum Monte Carlo calculations for stoquastic Hamiltonians do not suffer from the sign problem because the negativity of the nonzero off-diagonal matrix elements guarantees that the transition probabilities in the associated random walk are all positive.

In this article we first demonstrate that stoquastic Hamiltonians may be constructed which allow universal adiabatic quantum computation in a subspace. Then we show that the 3-local Hamiltonian problem is QMA-complete when restricted to stochastic Hamiltonians. These are Hamiltonians in which all matrix elements are real and non-negative, and the sum of matrix elements in any row or column is 1. Hence determining the lowest eigenstate of a symmetric stochastic matrix is QMA-hard. If $H$ is a stochastic Hamiltonian, then $-H$ is stoquastic. Thus, our result also shows that determination of the highest-lying eigenstate of a stoquastic matrix is QMA-hard, sharpening the intuition that it is the positivity of the ground state which causes its local Hamiltonian problem to fall into a classical class. We then show that universal
adiabatic quantum computation is possible in the ground state of a stochastic frustration-free Hamiltonian. Defining the computational problem stochastic $k$-SAT in analogy to the definition of stoquastic $k$-SAT given in [3], we show that this problem is QMA$_1$-complete for $k = 6$. (QMA$_1$ is a slight variant of QMA such that in YES instances, Arthur can be made to accept with probability one [16].)

II. QMA-COMPLETENESS AND ADIABATIC UNIVERSALITY OF STOQUASTIC HAMILTONIANS

We start with the result of [17], which shows that for a Hamiltonian of the form

$$H_{XZ} = ∑_i d_i X_i + ∑_i h_i Z_i + ∑_{i,j} K_{ij} X_i X_j + ∑_{i,j} J_{ij} Z_i Z_j,$$

(2)

the 2-local Hamiltonian problem is QMA-complete if the coefficients $d_i$, $h_i$, $K_{ij}$, and $J_{ij}$ are allowed to have both signs. Furthermore, time-dependent Hamiltonians that take the form $H_{XZ}$ at all times can perform universal adiabatic quantum computation [17].

Starting with a Hamiltonian of the form $H_{XZ}$ on $n$ qubits, we can eliminate the negative matrix elements in each term using a technique from [18]. Essentially, the idea is that instead of representing the group $T_k$ can to a large degree protect the entire computation. Note that an energy penalty against the ancilla qubit leaving the $|−⟩$ subspace exactly matches the spectrum of $H_{XZ}(t)$, the only difference being the addition of an ancilla qubit in the $|−⟩$ state. Because $\tilde{H}_{XZ}(t)$ has no coupling between the $|−⟩$ subspace and the $|+⟩$ subspace, the adiabatic theorem may be applied within the $|−⟩$ subspace. The relevant eigenvalue gap is thus the same as that of $H_{XZ}(t)$, and so is the run time.

In standard adiabatic quantum computation, the qubits are in the ground state of the instantaneous Hamiltonian. Thus, any disturbance to the state costs energy. This is thought to offer some protection against thermal noise [19]. When performing universal adiabatic quantum computation with $\tilde{H}_{XZ}(t)$, the qubits are not in the ground state. However, this can only occur by disturbing the ancilla qubit out of the state $|−⟩$. By protecting the ancilla qubit, one can to a large degree protect the entire computation. Note that an energy penalty against the ancilla qubit leaving the state $|−⟩$ would be nonstoquastic. This is why the preceding construction fails to prove QMA-completeness and universal adiabatic quantum computation using the ground state of a stoquastic Hamiltonian, as we expect it must, based on the complexity-theoretic results of [2–4,20].

III. QMA-COMPLETE PROBLEMS FOR MARKOV MATRICES

The second main result of our article provides an example of a QMA-complete classical problem: finding the lowest eigenvalue of a symmetric Markov matrix. A matrix with all non-negative entries, such that the entries in any given

$$\bar{H}_{XZ} = -∑_k α_k |T_k|,$$

(8)
column sum to 1 is called a stochastic or Markov matrix. These matrices are named after Markov chains, which are stochastic processes such that, given the present state, the future state is independent of the past states. Suppose a system has $d$ possible states. Then, its probability distribution at time $t$ is described by the $d$-dimensional vector $x_t$ whose entries are non-negative and sum to 1. If the system is evolving according to a Markov process, then its dynamics are completely specified by the equation $x_{t+1} = M x_t$, where $M$ is a $d \times d$ stochastic matrix.

Markov processes for which the Markov matrix is symmetric correspond to random walks on undirected weighted graphs. (Self-loops are allowed and correspond to diagonal matrix elements.) These matrices are doubly stochastic: The sum of the entries in any row or column is 1. By the Perron-Frobenius theorem, the highest eigenvalue of a symmetric stochastic matrix is 1, and the corresponding eigenvector is the uniform distribution. The eigenvalue with next largest magnitude corresponds to random walks on undirected weighted graphs. A symmetric stochastic matrix is Hermitian and therefore one can also think of these matrices as Hamiltonians. To prove that finding the lowest eigenvalue of a 3-local symmetric stochastic matrix is QMA-complete, we again use a reduction from the QMA-complete $H_{XZ}$ Hamiltonian of [17]. We must take the opposite sign convention from Eq. (3),

$$H_{XZ} = \sum_k \alpha_k S_k,$$  

where $S_k = -T_k$. Now define

$$\hat{H}_{XZ} = \frac{1}{N} \sum_k \alpha_k \bar{S}_k,$$  

where

$$N = \sum_k \alpha_k$$

and $\bar{S}_k$ is the permutation matrix obtained by applying the replacement rules (5) to $S_k$. By construction, $\hat{H}_{XZ}$ is a 3-local, symmetric, doubly stochastic matrix. We can rewrite $H_{XZ}$ as

$$\hat{H}_{XZ} = \frac{1}{N} \left( H_{XZ} \otimes |-| - |+\rangle \langle +| + H_{XZ} \otimes |+\rangle \langle +| + \right),$$

where $H_{XZ} = \sum_k \alpha_k |S_k|$. Thus, to determine an eigenvalue of $H_{XZ}$ to within $\pm \epsilon$ we must find the corresponding eigenvalue of $\hat{H}_{XZ}$ to within $\pm \epsilon / N$. Because $H_{XZ}$ is a 2-local Hamiltonian on $n$ qubits with coupling strengths of order unity, $N$ is at most $O(n^2)$. Thus, the problem of determining the eigenvalue of $H_{XZ}$ corresponding to the ground state of $H_{XZ}$ to polynomial precision is QMA-hard.

To obtain a cleaner QMA-hard problem, we would like to construct a stochastic matrix whose lowest eigenvalue is QMA-hard to find. To do this, let

$$H_p = (1-p)\sigma_{n+1}^+ + p \hat{H}_{XZ}.$$

Here $\sigma_{n+1}^+ = |+\rangle \langle +| = \frac{1}{2}(1 + X_{n+1})$ acts on the ancilla qubit, thereby giving it an energy penalty of size $(1-p)$ against leaving the state $|\rangle$. For $0 \leq p \leq 1$, $H_p$ is a stochastic Hamiltonian. For $p < 1/3$, the energy penalty is large enough that the highest eigenvalue in the $|\rangle$ subspace lies below the lowest eigenvalue in the $|+\rangle$ subspace. Thus, our problem is the adiabatic theorem,2 1 be a state of $H_{XZ}$ scaled by $p/N$, and the upper half of the spectrum of $H_p$ is the spectrum of $\hat{H}_{XZ}$ scaled by $p/N$ and shifted up by $1-p$.

Thus, we can obtain the ground energy of $H_{XZ}$ to polynomial precision by computing the lowest eigenvalue of $H_p$ to a higher but still polynomial precision. This reduction proves that finding the lowest eigenvalue of $H_p$ to polynomial precision is QMA-hard. Using the quantum algorithm for phase estimation, one easily shows that the problem of estimating the lowest eigenvalue of $H_p$ is contained in QMA (see [11]). Thus, this problem is QMA-complete.

IV. FRUSTRATION-FREE ADIABATIC COMPUTATION

It was stated in [3] that universal adiabatic quantum computation can be performed in the ground state of a 5-local frustration-free Hamiltonian. Let $U = U_1 \cdots U_2 U_1$ be a quantum circuit acting on $n$ qubits with $L = \text{poly}(n)$ gates. Let

$$|\psi_j\rangle = U_j \cdots U_1 |0\rangle^\otimes n$$

be a state of $n$ qubits corresponding to the $j$th state of the time evolution of a quantum circuit specified by gates $U_j$ and

$$|c_j\rangle = |1^{s+1}0^{L-s}\rangle.$$

be a state of $L + 1$ clock qubits. Bravyi and Terhal construct a parametrized 5-local Hamiltonian $H(s)$ such that the ground state $|\psi(s)\rangle$ satisfies

$$|\psi(1)\rangle = \frac{1}{\sqrt{L+1}} \sum_{j=0}^L |\psi_j\rangle |c_j\rangle,$$

$$|\psi(0)\rangle = \frac{1}{\sqrt{L+1}} \sum_{j=0}^L |0^s\rangle |c_j\rangle.$$  

We can think of the first register in $|\psi(s)\rangle$ as consisting of “work” qubits on which the computation happens and the second register in $|\psi(s)\rangle$ as being a clock containing a time written in unary.

For $s \in [0, 1]$, the minimal eigenvalue gap between the ground state and first excited state of $H(s)$ is $O(1/L^2)$. By the adiabatic theorem,2 1 be a polynomial eigenvalue gap ensures that given $|\psi_0\rangle$, one obtains $|\psi_1\rangle$ by applying $H(s)$ and varying $s$ from 0 to 1 over poly($L$) time. By measuring the clock register of $|\psi(1)\rangle$, one obtains the result $|1^{L+1}\rangle$ with probability $1/(L+1)$. If this result is obtained, one finds the output of the circuit $U$ by measuring the first register of qubits in the computational basis. By repeating this process with $O(L)$ copies of $|\psi(1)\rangle$, one succeeds with high

2Many versions of the adiabatic theorem have been proven. For one example, see Appendix F of [21].
probability. Alternatively, one can pad the underlying circuit with $L$ identity gates, in which case each trial succeeds with probability 1/2.

The construction from [3] invokes the fact that the spectrum of $H(1)$ is independent of the form of the gates $U_j$. By choosing a gate set which is composed of elements of simply connected unitary groups such as $SU(2)$ and $SU(4)$, one may construct a continuous path connecting each gate to the identity and use a single parameter $s$ to transform all gates from the identity to the final circuit at once. The Hamiltonian at $s = 0$ corresponds to the identity circuit, and its ground state is the uniform superposition of the clock states tensored with the initial data on the work qubits. In this ground state, the qubits of the clock register are entangled. It is standard to design adiabatic computations such that the initial Hamiltonian has a product state as its ground state, because such states should be easily produced by cooling or single-qubit measurements. In this section we construct a modified version of the construction from [3] that satisfies this condition and is still frustration free.

Let $c(j)$ indicate the $j$th clock qubit and let $w(j)$ indicate the $j$th work qubit. Let

$$H^\text{init}_j = |1\rangle\langle 1|_{w(j)} \otimes |10\rangle\langle 10|_{c(1),c(2)},$$

$$H^\text{clock}_j = |01\rangle\langle 01|_{c(j-1),c(j)}.$$ 

For $j \in \{1, \ldots, L - 1\}$, define

$$H^\text{prop}_j(s) = s|10\rangle\langle 10|_{c(j),c(j+1),c(j+2)} + (1-s)|01\rangle\langle 01|_{c(j),c(j+1),c(j+2)} - \sqrt{s(1-s)}U_j \otimes |10\rangle\langle 10|_{c(j),c(j+1),c(j+2)} + U_j^\dagger \otimes |01\rangle\langle 01|_{c(j),c(j+1),c(j+2)}$$

and let

$$H^\text{prop}_L(s) = s|10\rangle\langle 10|_{c(L),c(L+1)} + (1-s)|11\rangle\langle 11|_{c(L),c(L+1)} - \sqrt{s(1-s)}U_L \otimes |10\rangle\langle 10|_{c(L),c(L+1)} + U_L^\dagger \otimes |11\rangle\langle 11|_{c(L),c(L+1)}.$$

It can be directly verified that each $H^\text{clock}$, $H^\text{init}$, and $H^\text{prop}(s)$ is a projector. Here, for convenience, we define $s$ so that it varies from 0 to 1/2 rather than from 0 to 1 as is done in [3]. Our frustration-free Hamiltonian is the following sum of projectors:

$$H^\text{clock} = |0\rangle\langle 0|_{c(0)} + \sum_{j=1}^L H^\text{clock}_j,$$

$$H^\text{init} = \sum_{j=1}^n H^\text{init}_j,$$

$$H^\text{prop}(s) = \sum_{j=1}^L H^\text{prop}_j(s),$$

$$H^\text{FF}(s) = H^\text{clock} + H^\text{init} + H^\text{prop}(s).$$

If $U_1 \cdots U_L$ are chosen from a universal set of two-qubit gates, then $H^\text{FF}(s)$ is an efficient 5-local frustration-free adiabatic quantum computer. To see how this Hamiltonian achieves universal adiabatic computation, we examine the various terms one by one. The ground state of $H^\text{FF}(s)$ is the simultaneous zero eigenspace of $H^\text{clock}$, $H^\text{init}$, and $H^\text{prop}$. $H^\text{clock}$ commutes with $H^\text{FF}(s) + H^\text{init}$ and provides an energy penalty of at least unit size if the clock register is not in one of the unitary states $|c_j\rangle = |1^{j+1}0^{L-j}\rangle$. Thus, the low-lying spectrum of $H^\text{FF}(s)$ is strictly contained in the ground space of $H^\text{clock}$.

For any bit string $x \in \{0, 1\}^n$ and integer $j \in \{1, 2, \ldots, L\}$, let

$$|\chi_j^x\rangle = (U_jU_{j-1}\cdots U_1|x\rangle) \otimes |c_j\rangle,$$

where $|c_j\rangle$ is as defined in Eq. (13). (We also define $|\chi_j^0\rangle = |x\rangle \otimes |c_0\rangle$.) There are $2^n(L+1)$ such states and they form an orthonormal basis for the ground space of $H^\text{clock}$. In this basis, $H^\text{prop}(s) + H^\text{init}$ takes the block-diagonal form

$$H^\text{prop}(s) + H^\text{init} = \bigoplus_{x\in\{0,1\}^n} M_x,$$

where

$$M_x = \begin{bmatrix}
\begin{array}{ccc}
s + |x| & -b & -b \\
-b & 1 & -b \\
-b & 1 & -b \\
\end{array}
\end{bmatrix}
$$

is an $L + 1$ by $L + 1$ matrix and $b = \sqrt{s(1-s)}$. Here $|x|$ denotes the Hamming weight of the bit string $x$. The appearance of $|x|$ is the sole manifestation of $H^\text{init}$. The rest of the matrix elements all come from the “hopping” action of $H^\text{prop}(s)$.

$M_{00\ldots 0}$ has the unique ground state

$$N \sum_{j=0}^L r^j |\psi_j\rangle |c_j\rangle,$$

where $|\psi_j\rangle$ and $|c_j\rangle$ are as defined in Eqs. (12) and (13), $r = \sqrt{\frac{s}{1-s}}$, and $N$ is a normalization factor. This constitutes the ground state of $H^\text{FF}(s)$. The first excited state of $M_{00\ldots 0}$ has energy $1 - 2\sqrt{s(1-s)}\cos\left(\frac{\pi}{L+1}\right)$. Because of the direct sum structure of $H^\text{FF}(s)$, we can apply the adiabatic theorem directly to $M_{00\ldots 0}$. The run time of the adiabatic algorithm is thus determined by the gap between the ground and first excited states of $M_{00\ldots 0}$. This takes its minimum at $s = 1/2$, where it is equal to $1 - \cos\left(\frac{\pi}{2(L+1)}\right)$, which is $O(1/L^2)$. For questions of fault tolerance, it is also useful to know the eigenvalue gap between the ground and first excited states of the full Hamiltonian $H^\text{FF}(s)$. The first excited energy of $H^\text{FF}(s)$ is equal to the ground energy of $M_{10\ldots 0}$, which is $1 - 2\sqrt{s(1-s)}\cos\left(\frac{\pi}{2(L+1)}\right)$. Thus the minimum eigenvalue gap of $H^\text{FF}(s)$ occurs at $s = 1/2$ and is equal to $1 - \cos\left(\frac{\pi}{2(L+1)}\right)$, which is $O(1/L^2)$.

By Eq. (14), the ground state of $H^\text{FF}(0)$ is $|00\ldots 0\rangle \otimes |1000\ldots 0\rangle$, and the ground state of $H^\text{FF}(1/2)$ is the same state $\sqrt{\frac{1}{L+1}} \sum_{j=0}^L |\psi_j\rangle |c_j\rangle$ produced by the scheme of [3].
V. STOCHASTIC FRUSTRATION-FREE COMPUTATION

In [3], Bravyi and Terhal showed that adiabatic quantum computation in the ground state of a SFF Hamiltonian can be efficiently simulated by a classical computer. In this section we show that, in contrast, one can perform universal adiabatic quantum computation in the ground state of a stochastic frustration-free (StochFF) Hamiltonian \( H^{\text{StochFF}}(s) \).[Alternatively, we can view this as computation in the highest energy state of the stoquastic Hamiltonian \( -H^{\text{StochFF}}(s) \).]

It has been shown that the two-qubit controlled-NOT (CNOT) gate, together with any one-qubit rotation whose square is not basis preserving, are sufficient to perform universal quantum computation [22]. All matrix elements in these gates are real numbers. If we choose \( U_1, \ldots, U_L \) from this gate set, then \( H^{\text{FF}}(s) \) is a 5-local real frustration-free Hamiltonian.

Examining the construction of Sec. III, one sees that it can be applied to any Hamiltonian with real matrix elements, and it increases the locality by one. This construction also preserves frustration freeness, as we will show in the next paragraph. We can thus use this construction on \( H^{\text{FF}}(s) \) to obtain a 6-local stochastic frustration-free Hamiltonian whose ground state is universal for adiabatic quantum computation.

To show that the mapping of Sec. III preserves frustration freeness, consider applying this mapping to a frustration-free local Hamiltonian \( H = \sum_{j=1}^{N} H_j \), where \( H_j = \sum_a \alpha_j^a S_k^a \) (where each \( S_k^a \) is, up to an overall sign, a tensor product of Pauli operators and each \( \alpha_j^a \) is positive). We obtain the Hamiltonian

\[
H_p = p H + (1 - p) \left( \frac{I + X_{n+1}}{2} \right)
= \sum_j \frac{N_j}{N} \left[ p H_j + (1 - p) \left( \frac{I + X_{n+1}}{2} \right) \right],
\]

where \( N_j = \sum_a \alpha_j^a \) and \( N = \sum_j N_j \). When \( p < \frac{1}{2} \), \( H_p \) is stochastic and has a zero-energy ground state with an eigenvalue gap which is \( \frac{N}{2} \) times the gap of \( H \). Furthermore, we see from (15) (and the fact that each \( H_j \) is positive semidefinite) that \( H_p \) is a sum of positive semidefinite operators. Hence, the Hamiltonian \( H_p \) is frustration free.

VI. GENERALIZATIONS

The constructions of Secs. III and II replace Hamiltonians with real matrix elements of both signs by computationally equivalent Hamiltonians with real positive matrix elements. In this section we show that this technique can be generalized to directly replace Hamiltonians with complex matrix elements by computationally equivalent Hamiltonians with only real positive matrix elements. However, in the process we necessarily introduce a twofold degeneracy of the ground state.

Let \( H \) be an arbitrary \( k \)-local Hamiltonian. We may expand \( H \) as

\[
H = \sum_j \alpha_j O_j,
\]

where each \( O_j \) is a tensor product of \( k \) or fewer Pauli matrices and each \( \alpha_j \) is positive. Each entry in each \( O_j \) is \( \pm 1 \) or \( \pm i \).

We can replace the group \( \{ 1, i, -1, -i \} \) with its left-regular representation

\[
\begin{align*}
1 & \mapsto F, \\
-1 & \mapsto F^2, \\
-i & \mapsto F^3, \\
1 & \mapsto F^4,
\end{align*}
\]

where

\[
F = \begin{pmatrix}
0 & 1 & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1 \\
1 & 0 & 0 & 0
\end{pmatrix}.
\]

The eigenvectors of \( F \) are \( |v_0\rangle, |v_1\rangle, |v_2\rangle, |v_3\rangle \), where

\[
|v_j\rangle = \frac{1}{2} \sum_{i=0}^{3} \langle i | j \rangle.
\]

The corresponding eigenvalues are

\[
F|v_j\rangle = i^j |v_j\rangle.
\]

Let \( S_j \) and \( A_j \) be the real and imaginary parts of \( \alpha_j O_j \). That is, \( S_j \) and \( A_j \) are the unique real symmetric and antisymmetric matrices such that

\[
\alpha_j O_j = S_j + i A_j.
\]

Further, let \( S_j^+ = (S_j + S_j) / 2 \) and \( S_j^- = (S_j - S_j) / 2 \), and similarly for \( A_j^\pm \), where | \cdot | denotes the entrywise absolute value. Applying the replacement (17) to \( H \) and dividing by \( N = \sum_j \alpha_j \) yields the stochastic Hamiltonian \( \tilde{H} \) with the decomposition

\[
\tilde{H} = \frac{1}{N} \left( H^{(0)} \otimes |v_0\rangle\langle v_0| + H^{(1)} \otimes |v_1\rangle\langle v_1| + H^{(2)} \otimes |v_2\rangle\langle v_2| + H^{(3)} \otimes |v_3\rangle\langle v_3| \right),
\]

where

\[
H^{(0)} = \sum_j S_j^+ + S_j^- + A_j^+ + A_j^-,
\]

\[
H^{(1)} = \sum_j S_j^+ - S_j^- + i A_j^+ - i A_j^-,
\]

\[
H^{(2)} = \sum_j S_j^+ + S_j^- - A_j^+ - A_j^-,
\]

\[
H^{(3)} = \sum_j S_j^+ - S_j^- - i A_j^+ + i A_j^-.
\]

\( H^{(1)} = H \); thus, the spectrum of \( \tilde{H} \) in the \( |v_1\rangle \) subspace matches that of \( H \) up to a normalization factor of \( N \) and a pair of extra ancilla qubits. If we write each projector \( |v_j\rangle\langle v_j| \) in terms of the Pauli basis we obtain
\[
\begin{align*}
|v_0\rangle\langle v_0| &= \Pi_X^0 \Pi_Y^0, \\
|v_1\rangle\langle v_1| &= \Pi_X^0 \Pi_Y^+, \\
|v_2\rangle\langle v_2| &= \Pi_X^+ \Pi_Y^0, \\
|v_3\rangle\langle v_3| &= \Pi_X^+ \Pi_Y^+.
\end{align*}
\]
where $\Pi_k$ is the projector onto the eigenvalue $\pm 1$ eigenstate of the Pauli matrix $a$. Thus, an $X$ penalty on the first ancilla qubit will separate the $|v_1\rangle$, $|v_2\rangle$ subspace from the $|v_0\rangle$, $|v_2\rangle$ subspace. So, taking, $0 < p < \frac{1}{3}$, the stochastic Hamiltonian

$$H'_p = (1 - p) \left( \frac{I + X_{n+1}}{2} \right) + p \tilde{H}$$

has ground space spanned by $|\psi^{(1)}\rangle|v_1\rangle$ and $|\psi^{(3)}\rangle|v_2\rangle$, where $|\psi^{(1)}\rangle$ is the ground state of $H^{(1)}$ and $|\psi^{(3)}\rangle$ is the ground state of $H^{(3)}$. $H^{(1)} = H$; thus, $|\psi^{(1)}\rangle$ is the ground state of $H$. $H^{(3)} = H'$; thus, $|\psi^{(3)}\rangle$ is the complex conjugate of the ground state of $H$.

A simple argument shows that the doubling in the spectrum of $H'$ is a necessary property for any construction which maps an arbitrary Hamiltonian onto a real Hamiltonian $H_R$, where $H_R$ is equal to $H$ within a fixed 1D subspace of the ancillas. Suppose that we have such a map which sends an arbitrary Hamiltonian $H$ which acts on a Hilbert space $H_1$ to a real Hamiltonian $H_R$ on a larger Hilbert space $H_1 \otimes H_2$ with the property that

$$H_R = H \otimes \ket{\phi} \bra{\phi} + H^{\text{other}} \otimes (I - \ket{\phi} \bra{\phi}),$$

where the state $|\phi\rangle \in H_2$ does not depend on the particular Hamiltonian $H$ but the operator $H^{\text{other}}$ may depend on $H$. Then for any eigenvector $|\psi\rangle$ of $H$ with energy $E$, we have

$$H_R|\psi\rangle = E|\psi\rangle.$$ (21)

Since $H_R$ is real, complex conjugating this equation gives

$$H_R|\psi^*\rangle = E|\psi^*\rangle.$$ (22)

To show that doubling exists in the spectrum, it is sufficient to show that $\langle \phi|\phi^*\rangle = 0$. To prove this, first use Eq. (23) to obtain

$$(I \otimes \ket{\phi} \bra{\phi}) H_R|\psi^*\rangle = E|\psi^*\rangle \langle \phi|\phi^*\rangle.$$ (24)

Then use Eq. (21) to obtain

$$(I \otimes \ket{\phi} \bra{\phi}) H_R|\psi^*\rangle \langle \phi^*| = (H|\psi^*\rangle) \langle \phi|\phi^*\rangle.$$ (25)

Equating these expressions gives

$$H|\psi^*\rangle \langle \phi|\phi^*\rangle = E|\psi^*\rangle \langle \phi|\phi^*\rangle.$$ (26)

This must hold for all Hamiltonians $H$ and eigenstates $|\psi\rangle$, and therefore it must be the case that $\langle \phi|\phi^*\rangle = 0$. So we have shown that the doubling in the spectrum of $H'$ is a necessary feature of the type of maps we consider. For constructing universal adiabatic quantum computers, the degeneracy induced by this construction may be problematic. However, for proving complexity-theoretic completeness results, it is often irrelevant, as we see in the next section.

VII. STOCHASTIC $k$-SAT

The methods of the previous section can be used to show, roughly speaking, that deciding whether or not a Hamiltonian which is a sum of positive semidefinite stochastic operators is frustration free is as difficult as the general problem of deciding whether a Hamiltonian is frustration free. In this section we formalize this by defining a problem called stochastic $k$-SAT, which we show to be QMA$_1$-complete for $k = 6$.

We first recall the definition of stoquastic $k$-SAT which is given in [3].

Problem: Stoquastic $k$-SAT

Input. A set of $k$-local Hermitian operators $\{H_j\}$ for $j \in \{1, \ldots, m\}$, where $m = \text{poly}(n)$ and a parameter $\epsilon > 1/\text{poly}(n)$:

(i) each $H_j$ is positive semidefinite;

(ii) each $H_{j}$ has norm which is bounded by a polynomial in $n$;

(iii) every $H_j$ is stoquastic.

Output. If $H = \sum_j H_j$ has a zero-energy ground state, then this is a YES instance. Otherwise, if every eigenstate of $H$ has energy $> \epsilon$, then it is a NO instance.

Promise. Either the ground state of $H$ has energy 0, or else it has energy $> \epsilon$.

The stoquastic $k$-SAT problem is therefore the problem of deciding if a given stoquastic Hamiltonian that is a sum of positive definite operators is frustration free, given that either this is the case or else its ground energy exceeds $\epsilon$ [3]. Note that this definition of stoquastic $k$-SAT looks somewhat different from the definition of quantum $k$-SAT which was given in Sec. I, which was stated entirely in terms of projectors. Given an instance of stoquastic $k$-SAT, we can define operators $\Pi_j$ which project onto the zero eigenspaces of the $H_j$. When the Hamiltonians $H_j$ are stoquastic, these projectors are guaranteed to have non-negative matrix elements in the computational basis [3]. So given an instance of stoquastic $k$-SAT with Hermitian positive semidefinite operators $H_j$, it is possible to construct another instance of stoquastic $k$-SAT with operators $\tilde{H}_j = [1 - \Pi_j]$ that are all projectors.

We now define a problem called stochastic $k$-SAT, which is identical to stoquastic $k$-SAT except that condition (iii) is replaced by

(iii') Every $H_j$ is a stochastic matrix.

We note that there does not appear to be an equivalence between this definition of stochastic $k$-SAT and the corresponding definition where all the $H_j$ are (in addition) required to be projectors.

Given these two definitions and the foregoing map from an arbitrary Hamiltonian to a stochastic Hamiltonian, we now show how to reduce any instance of quantum 4-SAT to an instance of stochastic 6-SAT. Starting with an instance of quantum 4-SAT specified by a set of projectors $\{\Pi_j\}$ for $j \in \{1, \ldots, m\}$, we use the map of the previous section (with $p = \frac{1}{3}$ for concreteness) on each projector to obtain a set of 6-local positive semidefinite stochastic Hamiltonians $\{H_j\}$, where

$$H_j = \frac{2}{3} \left( \frac{I + X_{n+1}}{2} + \frac{1}{3} \Pi_j \right).$$ (27)

[Note that $\Pi_j$ refers to the operator obtained by applying the mapping from Eq. (20).] If the 4-SAT instance is satisfiable, then the stochastic 6-SAT instance will also be satisfiable. Define $N_{\text{max}}$ to be the maximum value of $N$ obtained for one of the terms $\Pi_j$ when using the mapping of Eq. (20). If the 4-SAT instance is not satisfiable, then for any state $|\phi\rangle$ there is some projector $\Pi_k$ such that $\langle \phi|\Pi_k|\phi\rangle \geq \frac{1}{m}$. If we take the parameter $\tilde{\epsilon}$ of the stochastic 6-SAT instance to be related to the parameter $\epsilon$ of the quantum 4-SAT instance
by \( \tilde{\epsilon} = \frac{a}{\log_2 n} \), then the stochastic 6-SAT instance will also be unsatisfiable. Therefore, stochastic 6-SAT is QMA_1-hard. Stochastic 6-SAT is contained in QMA_1 since every instance of stochastic 6-SAT can be mapped to an instance of quantum 6-SAT by taking projectors \( \Pi_j \) which project onto everything but the zero eigenspaces of the \( H_j \).

So QMA_1 completeness of stochastic 6-SAT follows from the results of Bravyi [16] on quantum k-SAT. This is in contrast to stoquastic k-SAT, which is contained in MA for every constant \( k \) [3].

VIII. QMA-COMPLETENESS FOR EXCITED STATES

The local Hamiltonian problem refers specifically to ground state energies. Similarly, we have formulated a computational problem based on the highest energy of a given Hamiltonian. It is natural to ask about the complexity of estimating the \( c \)th excited state. We can formulate this as follows. Let \( H \) be a \( k \)-local Hamiltonian on the Hilbert space \( \mathcal{H} \) of \( n \) qubits. Let \( \lambda_1 \leq \lambda_2 \leq \cdots \leq \lambda_{2^n} \) denote the eigenvalues of \( H \), corresponding to \( |\psi_1\rangle, |\psi_2\rangle, \ldots, |\psi_{2^n}\rangle \). The \( (k,c,\epsilon) \)-energy problem is as follows.

**Problem: \((k,c,\epsilon)\)-energy**

**Input.** We are given a classical description of \( f \), an integer \( c \geq 1 \), and a pair of parameters \( a, b \) such that \( b - a = \epsilon > 1/\text{poly}(n) \).

**Output.** If \( \lambda_c \leq a \), answer YES. If \( \lambda_c \geq b \), output NO.

**Promise.** \( H \) is such that the answer is YES or NO.

In this section we will show that the \( (k,c,\epsilon) \)-energy problem is QMA-complete for any \( c = O(1) \). Showing QMA-hardness is the easier of the two proofs. This can be achieved as follows. Let \( H^{(a)} \) and \( H^{(b)} \) be a pair of \( k \)-local Hamiltonians on \( n \) qubits, with spectra \( \lambda_1^{(a)}, \ldots, \lambda_{2^n}^{(a)}, |\psi_1^{(a)}\rangle, \ldots, |\psi_{2^n}^{(a)}\rangle \) and \( \lambda_1^{(b)}, \ldots, \lambda_{2^n}^{(b)}, |\psi_1^{(b)}\rangle, \ldots, |\psi_{2^n}^{(b)}\rangle \), respectively. Then

\[
H^{(ab)} = H^{(a)} \otimes |0\rangle\langle 0| + H^{(b)} \otimes |1\rangle\langle 1|
\]

is a \((k+1)\)-local Hamiltonian on \( n+1 \) qubits. Its complete set of eigenvalues is \( \lambda_1^{(a)}, \ldots, \lambda_{2^n}^{(a)}; \lambda_1^{(b)}, \ldots, \lambda_{2^n}^{(b)}, \) with corresponding eigenvectors \( |\psi_1^{(a)}\rangle|0\rangle, \ldots, |\psi_{2^n}^{(a)}\rangle|0\rangle; |\psi_1^{(b)}\rangle|1\rangle, \ldots, |\psi_{2^n}^{(b)}\rangle|1\rangle \). To prove QMA-hardness of a low-lying excited state, let \( H_0 \) be a Hamiltonian such that determining whether the ground energy is close to zero is QMA-hard. Given an integer \( c \), let \( d = \lfloor \log_2 c \rfloor \), \( P_k = \frac{1}{2} (Z_k + \mathbb{I}) \), and

\[
H_c = \sum_{k=0}^{d} 2^k P_k + \sum_{k=d+1}^{n} 2^{d+1} P_k - \left( c - \frac{1}{2} \right) \mathbb{I}.
\]

\( H_c \) has exactly \( c \) states with negative energy, and its lowest non-negative eigenvalue is \( \frac{1}{2} \). Thus determining the \( c \)th excited energy of \( H_c \otimes |0\rangle\langle 0| + H_0 \otimes |1\rangle\langle 1| \) is QMA-hard. In particular, it is interesting to note that by choosing \( c = 2 \) we construct a Hamiltonian whose eigenvalue gap between the ground state and the first excited state is QMA-hard to compute.

Next we show containment in QMA. The naive protocol would be for Merlin to provide Arthur with the state \(|\psi_1\rangle|\psi_2\rangle \ldots |\psi_c\rangle \) and for Arthur to use phase estimation to check that the \( c \) registers each contain a state of energy at most \( a \). The problem is that for NO instances there are many ways for Merlin to cheat. For example if \( \lambda_1 \leq a \) but \( \lambda_c \geq b \), the answer is NO, but Merlin can provide the state \( |\psi_1\rangle|\psi_2\rangle \ldots |\psi_c\rangle \) as a supposed witness. To prevent this, Arthur needs to somehow check that he has been given a set of \( c \) orthogonal states that each have energy at most \( a \). Thus, we propose the following protocol.

Arthur demands that Merlin give him the state

\[
|W\rangle = \frac{1}{\sqrt{c!}} \sum_{\pi \in S_c} \text{sgn}(\pi) |\psi_{\pi(1)}\rangle|\psi_{\pi(2)}\rangle \cdots |\psi_{\pi(c)}\rangle.
\]

Arthur performs the projective measurement to see that the state given to him by Merlin lies in the antisymmetric subspace of \( \mathcal{H}^{\otimes c} \). If this fails, he rejects. He then throws away all but the first register and performs phase estimation of \( H \) to precision better than \( \epsilon \). If the state has energy above \( b \), he rejects. Otherwise, he accepts.

It is clear that for YES instances, Arthur will accept the state \(|W\rangle\) with high probability. (The only source of error is imprecision in phase estimation.) We will next prove that for NO instances the acceptance probability is at most \( 1 - \frac{1}{c} \).

**Lemma 1.** For any state \(|\phi\rangle\) in the antisymmetric subspace of \( \mathcal{H}^{\otimes c} \) and any state \(|\alpha_1\rangle \in \mathcal{H}, (|\phi\rangle\langle\phi|)\otimes |1\rangle\langle 1|) |\phi\rangle \leq \frac{1}{c} |\phi\rangle \) where \( \otimes \) denotes the identity operator on \( \mathcal{H}^{\otimes (c-1)} \).

**Proof.** Extend \(|\alpha_1\rangle\) to an orthonormal basis \(|\alpha_1\rangle, |\alpha_2\rangle, \ldots, |\alpha_{2^n}\rangle \) for \( \mathcal{H} \). Let \( F \) be the set of functions \( f : \{1, 2, \ldots, c\} \to \{1, 2, \ldots, 2^n\} \) such that \( f(1) < f(2) < \cdots < f(c) \). Thus, \(|F| = \binom{c}{2}\). For any \( f \in F \) we have the following Slater determinant:

\[
|D_f\rangle = \frac{1}{\sqrt{c!}} \sum_{\pi \in S_c} \text{sgn}(\pi) |\alpha_{f(1)}\rangle|\alpha_{f(2)}\rangle \cdots |\alpha_{f(c)}\rangle.
\]

It is a standard result that these \( \binom{c}{2} \) states form a complete orthonormal basis for the antisymmetric subspace of \( \mathcal{H}^{\otimes c} \). For any \( f, g \in F \), we have

\[
\langle D_f | (|\alpha_1\rangle \otimes |1\rangle \otimes |D_g\rangle = \frac{\delta_{f,g} \delta_{f(1),1}}{c},
\]

where each \( \delta \) denotes a generalized Kronecker-\( \delta \). Because \(|\phi\rangle\) is antisymmetric, it can be decomposed in the Slater determinant basis,

\[
|\phi\rangle = \sum_{f \in F} \phi_f |D_f\rangle,
\]

and \( \phi_f = \phi_f^* \phi_f \) is a corresponding probability distribution on \( F \). Thus,

\[
\langle \phi | (|\alpha_1\rangle \otimes |1\rangle) |\phi\rangle = \sum_{f,g \in F} \phi_f^* \langle D_f | (|\alpha_1\rangle \otimes |1\rangle \otimes |D_g\rangle \phi_g.
\]

By Eq. (29), this is

\[
\frac{1}{c} \sum_{f \in F} \phi_f \delta_{f(1),1} \leq \frac{1}{c}.
\]

By Eq. (29), this is
The quantity
\[ p_j(\phi) = \langle \phi | (\psi_j)(\psi_j) \otimes \mathbb{I} | \phi \rangle \]
is the probability of obtaining |\psi_j\rangle if we measure the first register of a state |\phi\rangle in the eigenbasis of \( H \). By Lemma 1,
\[ \sum_{j=1}^{c-1} p_j(\phi) \leq 1 - \frac{1}{c}. \]
Thus, with probability at least \( \frac{1}{c} \), such a measurement would yield |\psi_j\rangle, with \( j > c - 1 \). Thus, if \( \lambda_c > b \), then with probability at least \( \frac{1}{c} \) a measurement of the observable \( H \) would yield energy of at least \( b \). The phase estimation algorithm can in poly(1/\( \epsilon \)) time perform such an energy measurement with an exponentially smaller chance of making an error as large as \( \epsilon \). Thus, the protocol is sound, which completes the proof that the \( (c, k, \epsilon) \)-energy problem is QMA-complete for constant \( c \) and \( k \) and polynomially small \( \epsilon \).

**IX. DISCUSSION AND CONCLUSIONS**

The results presented in this article have several applications. Although calculating the ground-state energy of stoquastic Hamiltonians appears easier than calculating the ground-state energy of generic Hamiltonians, our results suggest that calculating other eigenstates of stoquastic Hamiltonians remains hard. Because the wave functions of these states have amplitudes which are both positive and negative, the hardness of determining their energy supports the intuition that it is the positivity of the amplitudes which makes the ground-state problem for stoquastic Hamiltonians easier. An extreme distinction between stochastic and stoquastic Hamiltonians arises when the Hamiltonians are also frustration free. Although adiabatic evolution with SFF Hamiltonians is simulable in BPP [3], we have shown that adiabatic evolution in the ground state of a stochastic frustration-free Hamiltonian is universal.

Second, these results may be relevant for the physical implementation of quantum computers. The first proof of universality of adiabatic quantum computation used 5-local interactions [25]. Since then, the Hamiltonians have been brought into incrementally more physically feasible form by various techniques while retaining universality [13,26,27].

The universal Hamiltonian \( H_{XZ} \) of [17] is one outcome of this chain of reductions. Here we add one more step to this chain, obtaining universal stochastic and stoquastic Hamiltonians which resemble those arising in some systems of superconducting qubits [4]. The constructions given here are at least 3-local, and so would require the use of perturbative gadgets to implement in terms of physical 2-local interactions.

Finally, our results are of interest from a purely complexity-theoretic point of view. Stochastic matrices arise outside the context of quantum mechanics, in Markov chains. Our reduction shows that finding the lowest eigenvalue of a certain class of exponentially large but efficiently describable doubly stochastic matrices is QMA-complete. (These stochastic matrices correspond to Markov chains in which the “update rule” is a probabilistic selection over some set of updates which are local in the tensor product sense.) In general, the problem of finding eigenvalues of stochastic matrices is of interest because the eigenvalue of second-largest magnitude determines the mixing time of the corresponding Markov chain. There exist Markov chains in which the eigenvalue of second-largest magnitude is negative and is the lowest-lying eigenvalue. We hope that the demonstration of a QMA-complete problem arising in a classical setting will help shed further light on the class QMA itself.

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