Direct certification of a class of quantum simulations

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(Dated: December 14, 2017)

One of the main challenges in the field of quantum simulation and computation is to identify ways to certify the correct functioning of a device when a classical efficient simulation is not available. Important cases are situations in which one cannot classically calculate local expectation values of state preparations efficiently. In this work, we develop weak-membership formulations of the certification of ground state preparations. We provide a non-interactive protocol for certifying ground states of frustration-free Hamiltonians based on simple energy measurements of local Hamiltonian terms. This certification protocol can be applied to classically intractable analog quantum simulations: For example, using Feynman-Kitaev Hamiltonians, one can encode universal quantum computation in such ground states. Moreover, our certification protocol is applicable to ground states encodings of IQP circuits demonstration of quantum supremacy. These can be certified efficiently when the error is polynomially bounded.

I. INTRODUCTION

Quantum devices offer the promise to outperform classical machines computationally and to solve problems intractable in the classical domain. This idea is most prominent in the idea of a fully-fledged universal quantum computer that could—once physically realised for large system sizes—solve problems like factoring in polynomial time [1]. Indeed, much effort is dedicated to realising the building blocks from which such a universal quantum computer could be built. What is already available with present technology, however, are quantum simulators [2], machines able to simulate other quantum systems, based on an idea going back to Richard Feynman [3]. In particular, this is true for so-called analog quantum simulators that involve a very large number of quantum constituents whose time evolution is not discretised as is the case in digital simulations. Prominent architectures for analog simulation are based on cold atoms in optical lattices [4], trapped ions [5] or superconducting qubits [6]. As of yet, their computational power is far from clear, though. They are special purpose machines, in that one can achieve a tremendous degree of control, but only over some parameters: In this way, it is perfectly conceivable to prepare ground states of interacting Hamiltonians to probe effects such as high-$T_c$ superconductivity [7] or lattice gauge theories [8]. Equally well, properties of time evolution can be probed in dynamical quantum simulators [9, 10].

In all such approaches the problem of certification arises, providing an answer to the question: Is the device working precisely in the anticipated fashion? In case of a quantum computation that solves a problem in NP, one can of course in retrospect efficiently certify the correctness of the solution to the decision problem by classical means. This approach is insufficient in many instances, however. To start with, not all problems interesting to be solved on a quantum device are decision problems. A particularly pressing question for quantum simulations, is to find out whether one has achieved a desired ground state preparation accurately. Moreover, it would be desirable to also obtain information about intermediate steps of a quantum computation or simulation. If parameter regimes are available that allow for an efficient classical simulation—as is the case, e.g., in non-interacting physical systems—one can set benchmarks in these regimes. In the physics literature, it is surprisingly common to assume that the only way to certify the correctness of a quantum simulation is to find regimes in which a classical simulation is available.

In this work, we show the converse: Ground state preparations of frustration-free Hamiltonians as an example of a type of quantum simulation can be certified efficiently using local measurements only, while at the same time the actual outcome of the simulation or quantum computation cannot be predicted classically. Our certification protocol is surprisingly simple and can be applied in a range of paradigmatic settings including the certification of IQP circuits [11], the toric code [12], graph states [13], and even arbitrary quantum computations encoded in Feynman-Kitaev-type Hamiltonians [14–17]. With the rapid experimental advances in the aforementioned experimental platforms, the large-scale realization of these models has come into reach. Our protocol yields an experimentally viable method to certify their correctness using experimental techniques that are available now.

The central idea of this work is to combine fidelity estimation techniques from quantum state tomography [18] with computational models from quantum complexity theory [19–21]. Its simplicity is the main virtue of our protocol—both on a conceptual level and in terms of the experimental capacities required for its implementation. Our main theoretical insight— that certification may well be possible even when classical simulation is not available—underpins the potential of quantum simulators as well as the necessity of their certification, hence addressing a timely topic in the quest for identifying quantum simulators with superior computational power. An important part of the purpose of this work is to provide a link between the physics and computer science literature, hence contributing to the debate on how to reliably certify quantum simulators outperforming classical machines.

Technically, we extend a known fidelity bound for the
ground states of frustration-free Hamiltonians \[^{[18]}\] Formulating the problem of fidelity estimation for ground states of frustration-free Hamiltonians as a weak-membership certification protocol \[^{[19]}\] then allows us to address the problem of rigorously certifying such ground states given an estimate of their energy only. To demonstrate that in this very setting, computationally intractable problems can be solved, we make use of the framework of a physically plausible variant of a Feynman-Kitaev Hamiltonian construction \[^{[14]}\]. Thus, small technical improvements on known results allow us to show our main result as formalised in Propositions \[^{[1]}\] and \[^{[2]}\] and make it applicable to real-world experimental settings.

An intriguing application of our certification protocol arises from Ref. \[^{[11]}\]. In this work, the authors present a restricted (non-universal) family of commuting quantum circuits, where sampling from their output probability distribution with an error in trace distance of less than $1/192$ is provably hard for any classical computer up to a plausible complexity-theoretic conjecture, unless the Polynomial Hierarchy (a generalisation of $P \neq NP$) collapses. This circuit can be mapped to the ground state of a frustration-free local Hamiltonian, where our certification method can be applied. Let us first note that from energy measurements alone one can in general not guarantee to accept all states within a constant error. For such a guarantee a polynomial bound on the preparation error is required. However, also for a constant allowed error, if the certification accepts a state, then the state is indeed within the required error bound. In this sense, the combination of our methods with the results of Ref. \[^{[11]}\] yield an experimental proposal to demonstrate quantum supremacy \[^{[20]}\].

### A. Related work

In computer science, notions of verified quantum computation \[^{[13],[21],[23]}\] address the question of how the correct functioning of quantum machines can be checked. In such approaches, it is usually a small, well-characterised quantum system that is used to certify the functioning of a larger quantum machine making use of the idea of interactive proof systems involving several rounds of interaction between a restricted verifier and a powerful prover. In multi-round systems, the dynamics of a system can even be certified without the need for any quantum mechanical capacities \[^{[24],[25]}\]. In the same spirit, it is shown in Refs. \[^{[26],[28]}\] how trust in a quantum simulator can be amplified to certifying an untrusted simulator or to learning about the Hamiltonian of an unknown system. Very recently, a single-round scheme was devised \[^{[23]}\] that relies on preparing the ground state of a Feynman-Kitaev Hamiltonian. Here, the authors consider the general situation in which casting verification as a QMA problem does not directly lead to a verification protocol, since the task of the verifier “may well be comparable in complexity to independently implementing the circuit to be evaluated.” For this general case the authors devise a rather sophisticated protocol, which, to an extent, seems to be skew to the conceptual simplicity desired of a quantum simulator.

### B. Notions of quantum simulation

We start our discussion by emphasising what properties of quantum simulators we consider indispensable. Except from solving an interesting problem from the physics perspective, a quantum simulator should satisfy the following two criteria: it should be certifiable (reliable), and it should be able to solve a computationally hard problem that cannot efficiently be solved using classical means (proper) (cf. Ref. \[^{[29]}\]).

(i) A quantum simulator should be proper. We call a ground state quantum simulator (GS-QS) proper if using it one can solve a computational task that is believed to be intractable on a classical computer.

(ii) A quantum simulator should be reliable. We call a GS-QS reliable if there exists a weak-membership certification test (Def. \[^{[3]}\]) by which the actual preparation $\rho_p$ of the ground state can be certified.

### C. Definitions

To begin with, let us define the key concepts for what follows. In particular, these concepts are properties of Hamiltonians relevant for ground state quantum simulations considered here, related to being local, frustration-free and having a Hamiltonian gap.

**Definition 1** ($k$-local Hamiltonians). Let $\Lambda$ be a finite set of sites, and associate with each site $v \in \Lambda$ a finite-dimensional Hilbert space $\mathcal{H}_v$. A Hamiltonian $H = \sum_{\lambda \in \Lambda} h_{\lambda}$ defined on $\mathcal{H} = \bigotimes_{v \in \Lambda} \mathcal{H}_v$ is called ($k$-)local if each term $h_{\lambda}$ acts on at most $k = O(1)$ sites within $\lambda$, i.e., has finite support $\text{supp}(h_{\lambda}) \leq k$ independent of the system size $N = |\Lambda|$.

Throughout, we assume the local terms to be bounded in spectral norm, $\|h_{\lambda}\| \leq O(1)$ for all $\lambda$ independent of the system size. Most important will be the situation of encountering spatial locality, where the supports of the $h_{\lambda}$ are connected sets on a physical lattice such as a cubic lattice. A Hamiltonian $H$ is said to be (polynomially) gapped if the energy gap $\Delta$ between the ground state energy $E_0$ and the first excited eigenenergy $E_1$ satisfies $\Delta = \Delta(N) \propto 1/\text{poly}(N)$. An important case where $E_0$ can be easily obtained is the case of frustration-free Hamiltonians. Hence, we can set $E_0 = 0$ in the following definition.

**Definition 2** (Frustration-free Hamiltonians). A local Hamiltonian $H = \sum_{\lambda} h_{\lambda}$ with ground-state energy $E_0 = 0$ is called frustration-free (FF), if the projection $P_0$ onto the ground-state space of $H$ satisfies $HP_0 = 0$ and $h_{\lambda}P_0 = 0$ for all $\lambda$. 
II. CERTIFICATION OF PROPER GROUND STATE QUANTUM SIMULATIONS

We now turn to the actual certification scheme and the main insight of this work: Quantum systems that are described by spatially local, frustration-free and gapped Hamiltonians with unique ground state realise (i) reliable GS-QS some of which are even (ii) proper. In other words, in a realistic physical setting, namely, for systems described by spatially local frustration-free gapped Hamiltonians with unique ground state it is indeed possible to solve problems that are believed to be intractable classically. This insight is very much in the original spirit of quantum simulators presumably outperforming classical devices.

To prove that GS-QS with frustration-free and polynomially gapped Hamiltonians with unique ground state is reliable we use that the ground state of such Hamiltonians be certified efficiently by an energy measurement only \([13]\). Considering the case in which this energy measurement is performed by measuring local Hamiltonian terms \([14, 17]\), we formalise this in Proposition \([11]\). There, we provide bounds on the required number of measurements given a desired accuracy.

To demonstrate that in the same setting also computationally hard problems can presumably be solved, we make use of the fact that universal quantum computation can be performed by cooling to ground states of frustration-free Hamiltonians \([14, 30]\). To demonstrate this one needs to show that any quantum computation can be reduced to (local) measurements on that ground state. Specifically, Proposition \([2]\) states that adiabatic quantum computation can be performed efficiently using a polynomially gapped spatially local frustration-free Hamiltonian \(H_{ac}\). Since adiabatic computation \((ac)\) is universal \([31]\), any problem in BQP can be solved using the system described by \(H_{ac}\).

A. Weak-membership certification

We now define weak-membership quantum state certification. This is at the heart of our notion of certifying the correctness of a preparation. It captures precisely what it means to “have prepared a state to a given accuracy in the laboratory”. Is is conveniently conceptualised in terms of a game between a sceptic and restricted certifier Arthur and a powerful but untrusted quantum prover Merlin. Arthur has access to classical computation and spatially local measurements; Merlin is able to prepare arbitrary quantum states of large quantum systems. The input to the game is the classical description of the Hamiltonian \(H\) the ground state of which we want to prepare. Merlin prepares a number of independent and identical copies of a quantum state \(\rho_p\). Arthur’s goal is to certify that \(\rho_p\) is indeed the ground state of \(H\). In our setting, the only interaction we require between Arthur and Merlin is the local measurements of Arthur on the state preparation given to him by Merlin \([19]\). The test accepts if the state \(\rho_p\) prepared by Merlin is sufficiently close to the actual ground state \(\rho_0\) in fidelity that is given by \(F(\sigma, \rho) := \text{Tr}[\sigma\rho]\) for at least one of \(\sigma\) and \(\rho\) pure. The fidelity is related to the trace distance \(d(\sigma, \rho) = \text{Tr}[|\sigma - \rho|]/2\) via \(1 - F(\sigma, \rho)^{1/2} \leq d(\sigma, \rho) \leq (1 - F(\sigma, \rho))^{1/2}\). A robust reading of what it precisely means for Arthur to “certify” \(\rho_p\) as a good approximation of \(\rho_0\) in terms of the fidelity is the notion of weak-membership state certification, which we illustrate in Fig. \([1]\).

Definition 3 (Weak-membership quantum state certification). Let \(F_T > 0\) be a threshold fidelity and \(0 < \alpha < 1\) a maximal failure probability. A test which takes as input a classical description of \(\rho_0\) and copies of a preparation of \(\rho_p\), and outputs “reject” or “accept” is a weak-membership certification test if with high probability \(1 - \alpha\) it rejects every \(\rho_p\) for which \(F(\rho_p, \rho_0) < F_T\), and accepts every \(\rho_p\) for which \(F(\rho_p, \rho_0) \geq F_T + \delta\) for some fidelity gap \(\delta > 0\).

B. Certification protocol

In this section, we lay out the actual certification protocol of frustration-free Hamiltonians. The anticipated state to be certified is the ground state \(\rho_0\) of \(H\), on which a sequence of local measurements is performed, followed by efficient classical post-processing.

Protocol 1 (Certification protocol for FF Hamiltonians).

1. Arthur chooses a threshold fidelity \(F_T < 1\), maximal failure probability \(1 > \alpha > 0\) and an estimation error \(\epsilon < (1 - F_T)/2\).

2. Now Arthur asks Merlin to prepare a sufficient number of copies of the ground state \(\rho_0\) of the frustration-free and gapped Hamiltonian \(H = \sum \lambda h_\lambda\), i.e., the unique state that fulfils \(H\rho_0 = 0\).
3. Arthur measures each of the $n$ local terms $m$ times on the copies of the state $\rho_p$ prepared by Merlin to determine an estimate $E^*$ of the expectation value $\sum_{\lambda} \text{Tr}[\rho_p h_{\lambda}]$. Each measurement is performed on a new copy, and $m$ is given by expression (5).

4. From the estimate $E^*$ he obtains an estimate $F_{\text{min}}^*$ of the lower bound $F_{\text{min}}$ on the fidelity $F = F(\rho_p, \rho_0)$ such that $F^*_{\text{min}} \in [F_{\text{min}} - \epsilon, F_{\text{min}} + \epsilon]$ with probability at least $1 - \alpha$.

5. If $F_{\text{min}}^* < F_T + \epsilon$ he rejects, otherwise he accepts.

The subsequent proposition makes the resources needed for such a certification more precise, and sets the fidelity gap $\Delta$ for which Protocol I is a weak-membership test.

**Proposition 1** (Weak-membership certification). Let $H = \sum_{\lambda} h_{\lambda}$ be a gapped Hamiltonian with unique ground state, ground state energy $E_0 := 0$, gap $\Delta$, and interaction strength $J = \max_{l} ||h_{\lambda}||$. Then Protocol I is a weak-membership certification test with fidelity gap $\delta = (1 - F_T) \left( 1 - \frac{\Delta}{\|H\|} \right) + \frac{2 \epsilon \Delta}{\|H\|}$, and requires

$$m \geq \frac{J^2 n^2}{2 \Delta^2 \epsilon^2} \ln \left( \frac{n + 1}{\ln(1 - 1/\alpha)} \right),$$

measurements of each of the $n$ local terms on $\rho_p$ to determine the expectation value $\langle H \rangle_{\rho_p} = \text{Tr}[H \rho_p]$.

With protocol I one is therefore able to efficiently certify ground state preparations of polynomially gapped Hamiltonians $H$ that are at least $1/\text{poly}(n)$ close to the target state in fidelity.

To keep the notation simple, we have assumed $E_0 = 0$. However, the general case with arbitrary $E_0$ can be reduced to it whenever the value of $E_0$ is known apriori. For instance, this is the case whenever $H$ is frustration-free, since the global ground state energy can be obtained from the local ones.

Moreover, in order to obtain the estimate $F_{\text{min}}^*$ of the fidelity lower bound $F_{\text{min}}$ the value of the gap $\Delta$ needs to be known.

Note that no assumptions on the prepared state $\rho_p$ are made, in particular, it need not be pure.

Note also that it does not matter how the measurement of the total energy $\langle H \rangle$ is performed; depending on the setup at hand, it may be more suitable to measure the energy directly rather than measure all terms individually as insinuated in Step 3 of the protocol. One major advantage of our approach is that one can perform the output measurement on the same copies as the certification measurement. This means that our certification protocol can be simply integrated in the read-out protocol of a GS-QS: perform the certification measurements on the copies of $\rho_p$ first and then use the same states for the readout measurements if the certification test accepts $\rho_p$.

### C. Proof of Proposition I

In order to show Proposition I we require some detail on how to estimate the global energy from measurements of local Hamiltonian terms, cast into the form of a large-deviation bound. Such a bound is given by the following Lemma, which is stated and proved along the same lines as Lemma S4 in Ref. [19].

**Lemma 1** (Estimation of the energy). Decompose the local terms in their eigenbasis: $h_{\lambda} = \sum_{\mu} e_{\lambda,\mu} P_{\lambda,\mu}$, where the $P_{\lambda,\mu}$ are orthogonal projections onto the eigenspaces corresponding to eigenvalues $e_{\lambda,\mu}$. Let $X^{(i)}_{\lambda}$ be the random variable that takes the value $e_{\lambda,\mu}$ with probability $\text{Tr}(\rho_p P_{\lambda,\mu})$ by the measurement of $h_{\lambda}$ on the $i^{\text{th}}$ copy of $\rho_p$. Moreover, let

$$\langle h_{\lambda} \rangle_{\rho_p}^* = \frac{1}{m} \sum_{i=1}^{m} X^{(i)}_{\lambda}$$

be the estimate of $\langle h_{\lambda} \rangle_{\rho_p}$ by a finite-sample average of $m$ measurement outcomes, and $\langle H \rangle_{\rho_p}^* = \sum_{\lambda} \langle h_{\lambda} \rangle_{\rho_p}^*$, the resulting estimate of $\langle H \rangle_{\rho_p}$. As above, define $J = \max_{l} ||h_{\lambda}||$. For $\tilde{\alpha} \geq 1/2$ it holds that

$$\mathbb{P} \left[ \left| \langle H \rangle_{\rho_p}^* - \langle H \rangle_{\rho_p} \right| \leq \epsilon \right] \geq \tilde{\alpha},$$

whenever

$$m \geq \frac{J^2 n^2}{2 \epsilon^2} \ln \left( \frac{n + 1}{\ln(1 / \tilde{\alpha})} \right).$$

**Proof.** By Hoeffding’s inequality,

$$\forall \lambda \in [n]: \mathbb{P} \left[ \left| \langle h_{\lambda} \rangle_{\rho_p}^* - \langle h_{\lambda} \rangle_{\rho_p} \right| \geq \epsilon \right] \leq 2 e^{-2 \epsilon^2 / m ||h_{\lambda}||^2},$$

since the $\{X^{(i)}_{\lambda}\}_\lambda$ are independent random variables and $0 \leq X^{(i)}_{\lambda} \leq ||h_{\lambda}||$. Eq. (6) is equivalent to

$$\forall \lambda \in [n]: \mathbb{P} \left[ \left| \langle h_{\lambda} \rangle_{\rho_p}^* - \langle h_{\lambda} \rangle_{\rho_p} \right| \leq \epsilon \right] \geq 1 - 2 e^{-2 m \epsilon^2 / ||h_{\lambda}||^2},$$

and since all measurements are independent

$$\mathbb{P} \left[ \forall \lambda \in [n]: \left| \langle h_{\lambda} \rangle_{\rho_p}^* - \langle h_{\lambda} \rangle_{\rho_p} \right| \leq \epsilon \right] \geq \prod_{\lambda} \left( 1 - 2 e^{-2 m \epsilon^2 / ||h_{\lambda}||^2} \right) \geq \left( 1 - 2 e^{-2 m \epsilon^2 / n J^2} \right)^n \geq \tilde{\alpha},$$

resulting in

$$\mathbb{P} \left[ \left| \langle H \rangle_{\rho_p}^* - \langle H \rangle_{\rho_p} \right| \leq \epsilon \right] \geq \left( 1 - 2 e^{-2 m \epsilon^2 / n J^2} \right)^n \geq \tilde{\alpha} .$$

Eq. (10) holds whenever

$$m \geq \frac{J^2 n^2}{2 \epsilon^2} \ln \left( \frac{2}{1 - \tilde{\alpha}^1/n} \right) =: m_{\text{opt}},$$

(11)
where we can upper-bound $m_{\text{opt}}$ as $m_{\text{opt}} \leq (J^2 n^2 / 2c^2) \ln \left( (n + 1) / \ln(1/\alpha) \right)$ [19]. This shows the claim.

We would like to bound the error between the actual fidelity $F(\rho_0, \rho_p)$ and the estimate thereof given the energy measurement $\langle H \rangle_{\rho_p}^*$. To that end, we first upper- and lower-bound the fidelity by the expectation value $\langle H \rangle_{\rho_p}$, thereby complementing the lower bound from Ref. [18].

**Lemma 2 (Bounds on the fidelity [18]).** Given a Hamiltonian $H$ with gap $\Delta > 0$ above the unique ground state $\rho_0$ with energy $E_0 = 0$, and maximum energy $E_{\text{max}} = \|H\|$. Suppose $\langle H \rangle_{\rho} < E_1$, the energy of the first excited state, for some prepared state $\rho$. Then the overlap $F(\rho_0, \rho)$ with the ground state is bounded as

$$F_{\max}(\rho) := 1 - \frac{\langle H \rangle_{\rho}^*}{\|H\|} \geq F(\rho_0, \rho) \geq 1 - \frac{\langle H \rangle_{\rho}}{\Delta} =: F_{\min}(\rho). \quad (12)$$

**Proof.** Decompose $H = \sum_{i=0}^n E_i |E_i\rangle \langle E_i|$. Then,

$$\|H\|(1 - \text{Tr}[\rho_0 \rho]) = E_{\max} \sum_{i=0}^n \langle E_i, \rho E_i \rangle \geq \text{Tr}[H \rho] \geq \Delta \sum_{i=0}^n \langle E_i, \rho E_i \rangle = \Delta (1 - \text{Tr}[\rho_0 \rho]), \quad (13)$$

( equivalently $\|H\|(1 - P_0) \geq H \geq \Delta (1 - P_0)$, where $P_0$ projects onto the ground space) yielding $1 - \text{Tr}[H \rho] / \|H\| \geq F(\rho_0, \rho) \geq 1 - \text{Tr}[H \rho] / \Delta$.

This means that the lower bound to the fidelity resulting from those estimates $F_{\min} := F_{\min}(\rho_p)^* = 1 - \langle H \rangle_{\rho_p}^* / \Delta$ obeys

$$\mathbb{P}[F_{\min}^* - F_{\min}^* \leq \epsilon] \geq 1 - \alpha, \quad (14)$$

with $F_{\min} = F_{\min}(\rho_p)$ whenever

$$m \geq \frac{J^2 n^2}{2\Delta^2 \epsilon^2} \ln \left( \frac{n + 1}{\ln(1/\alpha)} \right). \quad (15)$$

**Proof of Proposition 1** (i) (Completeness) Let $\rho_p$ be such that $F \geq F_T + \delta$ with $\delta$ given in Eq. [1]. Then

$$F \geq F_T + (1 - F_T) \left( 1 - \frac{\Delta}{\|H\|} \right) + \frac{2\epsilon \Delta}{\|H\|}, \quad (16)$$

which is equivalent to

$$F_T + 2\epsilon \leq 1 - \|H\|(1 - F) / \Delta. \quad (17)$$

Finally, it follows from Eq. [14] that $\mathbb{P}[F_{\min}^* - F_{\min}^* \leq \epsilon] \geq 1 - \alpha$, so that

$$\mathbb{P}[F_{\min}^* \geq F_T + \epsilon] \geq 1 - \alpha . \quad (18)$$

Hence, with probability larger than $1 - \alpha$ the test described in protocol [1] accepts $\rho_p$.

(ii) (Soundness) Let $\rho_p$ be such that $F \leq F_T$. It then follows that $F_{\min} \leq F < F_T$. Hence,

$$\mathbb{P}[F_{\min}^* < F_T + \epsilon] > 1 - \alpha \quad (19)$$

is implied by Eq. [14] so that $\rho_p$ is rejected with probability at least $1 - \alpha$.

**D. Use of the bounds in analog simulation**

The above techniques can readily be applied to frustration-free analog GS-QS. Indeed, many interesting Hamiltonians can be identified as frustration-free Hamiltonians in the above sense. Notably, for one-dimensional quantum systems, the parent-Hamiltonians of matrix-product states are frustration-free [32]. For two-dimensional systems, the toric code Hamiltonian [12] constitutes an important example of a Hamiltonian of this type.

This motivates the question of how such ground states of frustration-free Hamiltonians can be efficiently prepared. In the next section we discuss the preparation of ground states using an *adiabatic algorithm*. Depending on the setting other methods may prove more practical, though. For example, one can prepare the ground state by local unitary circuit or by engineering dissipation. In both of these cases, however, it is not yet fully known how to efficiently prepare ground states, unless additional conditions are satisfied. Specifically, if all local Hamiltonian terms commute, the ground state of a frustration-free Hamiltonian can be prepared efficiently using a unitary variant [33] of Moser’s algorithm for solving certain classical satisfiability problems [34]. Relatively, in Ref. [35] the authors demonstrate that some frustration-free Hamiltonians can be cooled into their ground states by engineering dissipation. In this scheme, the (possibly regrouped) local terms are individually cooled by engineering local dissipative terms suitably. If cooling one term cannot create an excitation at a different location the protocol is efficient.

**E. Encoding quantum computations in ground states**

Protocol [1] offers a convenient way to certify ground states of frustration-free Hamiltonians. To serve as proper quantum simulators, the problems solvable by means of these systems should be computationally hard, i.e., intractable on a classical computer. This is the case; indeed, one can perform universal (adiabatic) quantum computation by preparing the ground states of certain spatially local frustration-free Hamiltonian.
Proposition 2 (Encoding quantum computation in ground states [14]). Let $U = U_L \cdots U_1$ be an arbitrary quantum circuit involving one- and two-qubit unitaries $U_i$, $i = 1, \ldots, L$, that acts on $K$ qubits, and $L = \text{poly}(K)$, let $|\phi_0\rangle \in (C^2)^\otimes K$ be an initial state, and set $n = O(K^2)$.

Then there exists a Hamiltonian $H_{ac}(\lambda)$, $0 \leq \lambda \leq 1$ on an $n \times n$ square lattice each site of which can be occupied by either zero particles or one spin-$1/2$ particle. For the computation, a string of $2n$ particles move through the lattice. In particular, the following holds: For all $\lambda \in [0, 1]$, $H_{ac}(\lambda)$ is polynomially gapped as $\Omega(n^{-3})$. Moreover, the evolved Hamiltonian $H_{ac}(1)$ is frustration free and has a unique ground state $|\text{gs}_{ac}\rangle$. The probability of measuring the position of the particles on $|\text{gs}_{ac}\rangle$ such that one obtains the output state $(U_L \cdots U_1)|\phi_0\rangle$ in their spin degrees of freedom is lower bounded by a constant.

In the proof of Ref. [14] $H_{ac}$ is constructed as a variant of the Feynman-Kitaev Hamiltonian. Specifically, the Hamiltonian $H_{ac}$ describes the diffusion of a string of the particles on the lattice, where at each plaquette two particles interact via a four-body interaction term. That is, the position of the particle string serves as a clock register, their internal state as the work register. Proposition 2 entails that (a) the ground state of the Hamiltonian $H_{ac}(1)$ is unique, (2) can be efficiently prepared by preparing the ground state of $H_{ac}(0)$ and then tuning $\lambda$ adiabatically from 0 to 1, and (c) is polynomially gapped. Finally, (d) the output of the computation can be obtained by performing measurements on the spin degrees of freedom. Thus, the ground state of $H_{ac}(1)$ can be certified by an energy measurement on $|\text{gs}_{ac}\rangle$ according to Protocol 1. More recently, it has been shown how to improve this construction to make only use of nearest-neighbour interactions [34].

In order to familiarise the reader with the construction, here we sketch the most basic variant [30] of the Feynman-Kitaev clock construction [15-17]. In this construction, the discrete evolution under a quantum circuit is mapped to measuring the ground state of a local frustration-free Hamiltonian called the Feynman-Kitaev Hamiltonian. The discretised time evolution $(U_L \cdots U_2 U_1)|\phi_0\rangle_w$ of a $k$-qubit input state vector $|\phi_0\rangle_w$ is in this setting mapped onto a so-called history state

$$|\psi_{\text{hist}}\rangle = \frac{1}{\sqrt{L+1}} \sum_{i=0}^{L} (U_L \cdots U_0 |\phi_0\rangle_w) \otimes |t\rangle_c,$$

of a bipartite system comprising a work and a clock register. The work register consists of $k$ qubits, the clock-register is an $L$-dimensional quantum system with state vectors $\{|t\rangle\}_{t=0}^{L-1}$. The combined system is equipped with a Hilbert space $H_{\text{work}} \otimes H_{\text{clock}} \cong (C^2)^\otimes K \otimes C^L$ [37]. The history state is the ground state of a Hamiltonian that selects configurations for which the work register is updated correctly as the clock proceeds in time. Such a Hamiltonian has the form

$$H_{\text{update}} = \sum_{t=1}^{L} U_t \otimes |t\rangle \langle t - 1|_c + U_t^\dagger \otimes |t - 1\rangle \langle t|_c.$$

Whenever the clock undergoes a transition from $|t\rangle$ to $|t+1\rangle$, the unitary map $U_t$ is applied to the work register. The computation $U_t$ is undone via the application of $U_t^\dagger$ as the clock transits from $|t+1\rangle$ to $|t\rangle$. This Hamiltonian is of the form of a quantum walk on the line of state vectors $|\psi_t\rangle = (U_t \cdots U_0 |\phi_0\rangle_w) \otimes |t\rangle_c$. In each state vector $|\psi_t\rangle$ the computation up to step $t$ is stored in the work register as well as a time-marker in the clock register. The full Hamiltonian consists of an additional term $H_{\text{input}}$ that select the correct input configuration. Moreover, the highly non-local clock terms can be made strictly local using the construction of a unary clock in which the state vector $|t\rangle$ is represented by an $L$-qubit state vector

$$|t'\rangle = |1, \ldots, 1, 0, \ldots, 0\rangle.$$

The correct configurations of such a clock are enforced by another additional term $H_{\text{clock}}$. The full Feynman-Kitaev Hamiltonian then reads

$$H_{\text{FK}} = H_{\text{input}} + H_{\text{update}} + H_{\text{clock}}.$$  

The output of the computation can be obtained by measuring the clock register first, and then measuring the work register [38]. In order to raise the probability of obtaining outcome $L$ upon the clock measurement from $1/(L+1)$ to a constant, one can simply introduce another $O(L)$ identity gates after the last step of the computation.

The Hamiltonian [23] is gapped as $\Delta \propto 1/L^2$, it is 5-local (but not spatially local), and frustration-free with vanishing ground state energy, since all terms can be chosen as projections. The ground state can be prepared, for example, by adiabatically turning on the updating term $H_{\text{update}}$ [30]. If the preparation procedure starts out in the ground space manifold, and the adiabatic path is sufficiently smooth the adiabatic theorem tells us that eventually one ends up in the ground state of the final Hamiltonian $H_{\text{update}}$ in a time that scales as $1/\Delta^3$ [39].

Finally, note that Proposition 2 implies that arbitrary unitary and dissipative time-evolution can be simulated efficiently using the ground states of frustration-free and gapped Hamiltonians on a lattice. This is the case because arbitrary local Lindbladian dynamics can be approximated by a polynomial quantum circuit and thus simulated efficiently on a quantum computer [40].

III. TOWARDS QUANTUM SUPREMACY

In the previous section we have discussed a general mapping of quantum circuits to unique ground states of frustration-free, poly-gapped local Hamiltonians. In this section we will specialise to the family of circuits proposed by Bremner, Montanaro, and Shepherd [11], for which there is strong complexity-theoretic evidence (in contrast to Shor’s algorithm) that their output probability distribution is hard to
simulate on any classical computer within an error of $1/192$ in total variation distance. While this result on its own is remarkable, one might question whether the “theoretically hard” quantum state has actually been prepared in an experiment. However, using our certification result, one can indeed certify the sufficiently precise preparation of a quantum state, such that measurements in the $Z$-basis are intractable for classical computers under a plausible complexity-theoretic conjecture. This significantly strengthens the perspective to experimentally demonstrate quantum supremacy [20].

The result of Ref. [11] is based on the conjectured average-case hardness to compute the gap of degree-3 polynomials over $\mathbb{F}_2$, $f : \{0,1\}^n \to \{0,1\}$, which can be written (up to an additive constant) as

$$f(x) = \sum_{i,j,k} \alpha_{i,j,k} x_i x_j x_k + \sum_{i,j} \beta_{i,j} x_i x_j + \sum_i \gamma_i x_i \pmod{2},$$

where $\alpha_{i,j,k}, \beta_{i,j}, \gamma_i \in \{0,1\}$ and the gap is defined as $\text{gap}(f) = |\{x|f(x) = 0\}| - |\{x|f(x) = 1\}|$. Let the normalised gap be $\text{ngap}(f) = \text{gap}(f) / 2^n$. Then the following is assumed to be true.

**Conjecture 1** [11]. Let $f : \{0,1\}^n \to \{0,1\}$ be a uniformly random degree-3 polynomial over $\mathbb{F}_2$. Then it is $\#P$-hard to approximate $\text{ngap}(f)^2$ up to multiplicative error of $1/4 + o(1)$ for a $1/24$ fraction of polynomials $f$.

While Conjecture [11] is not known to hold for an uniformly random choice of polynomials, it is known to be true in the worst case [41]. All that is left to show is how to lift this result from worst-case to average-case hardness.

The gap as defined above can be expressed conveniently as the acceptance probability of a family $C_f$ of so-called IQP circuits on $n$ qubits, which has the following structure: (i) apply Hadamard gates on all qubits, (ii) apply a sequence of $Z$, $CZ$, $CCZ$ gates (encoding the terms of $f(x)$), (iii) apply Hadamard gates on all qubits again. That is, $\text{ngap}(f) = \langle 0 \otimes C_f | 0 \otimes \rho \rangle$. Note for experimental simplicity, that all the diagonal gates commute and the circuit has only constant depth.

We are now ready to state the result of Bremner, Montanaro, and Shepherd precisely:

**Theorem 1** [11]. Assume Conjecture [11] is true. If it is possible to classically sample from the output probability distribution of any IQP circuit $C$ in polynomial time, up to an error of $1/192$ in $\ell_1$ norm, then there is a $\text{BPP}^{NP}$ algorithm to solve any problem in $\text{P}^\#P$. Hence the Polynomial Hierarchy would collapse to its third level.

We would like to note that the Polynomial Hierarchy is of central importance in computational complexity theory as a generalisation of the better-known $P \neq NP$ conjecture.

**Theorem 1** requires to prepare the output distribution of an IQP circuit with error of $1/192$ in $\ell_1$ norm. Using our methods, quantum states closer than $O(1/poly(n))$ in terms of the system size $n$ can be certified. Thus sufficiently well-prepared states can be certified to be within the provably hard regime of quantum states that are hard to sample from.

Without further ado, let us now present the certification procedure proposed. Let $L$ be the number of gates in an IQP circuit.

1. Map the IQP circuit to the ground state $\rho_0$ of a frustration-free, gapped, local Hamiltonian, e.g., according to Proposition [2] and choose $m$ according to Eq. [5].

2. Flip a fair coin to decide for either (a) or (b):

   (a) Prepare the ground state $\rho_p$ of this local Hamiltonian adiabatically $m$ times, measure local Hamiltonian terms to certify an upper bound on $\|\rho_0 - \rho_p\|_1 \leq \varepsilon$, or

   (b) Prepare the ground state $\rho_p$ and measure the position of all spins which projects on the output state of the IQP circuit $\rho_1$ with constant probability (otherwise, repeat the step), then measure all qubits in the $Z$ basis.

Step [25] certifies, that the quantum simulator indeed prepares a state $\rho_p$ such that $\|\rho_0 - \rho_p\|_1 \leq \varepsilon(m)$. The ideal ground state $\rho_0$ of the Hamiltonian described in Proposition [2] has constant overlap with the output state of the IQP circuit $\rho_1$, i.e., $\|\rho_0 - \rho_1\|_1 \leq c$, which can therefore be project with constant probability on the subspace where the computation has completed by measuring the respective terms of the Hamiltonian, i.e.,

$$\rho_1 = \frac{\Pi_0 \Pi}{\text{Tr}(\Pi_0 \Pi)}.$$  

Similarly, we project in step [25] the physically prepared state

$$\rho_{L,p} = \frac{\Pi_0 \Pi}{\text{Tr}(\Pi_0 \Pi)}.$$  

It follows by the triangle inequality and properties of the trace distance, that

$$\|\rho_1 - \rho_{L,p}\|_1 \leq \frac{2}{c} \|\rho_0 - \rho_p\|_1 \leq \frac{2}{c} \varepsilon(m),$$

which can be driven below the required constant of $1/192$ by an appropriate choice of $m$. Thus, we can clearly certify that a state has been prepared with sufficient precision (i.e., within $1/192$ in total variation distance [42]) such that measuring the first $n$ qubits in the $Z$ basis results in a probability distribution from which it is hard to sample from classically by Theorem [1].

Since our ground state certification protocol only applies to states prepared to at least $O(1/poly(n))$ precision, we note that the protocol proposed above in fact asks for the preparation of states with higher quality (inverse polynomial accuracy) than what would be strictly necessary to satisfy the hardness result (constant accuracy). Nevertheless, we would
like to stress the experimental simplicity to implement reliable local measurements as compared to more complicated alternatives.

For example, the experimenter could as well choose to implement a constant-accuracy fidelity estimation protocol using Hamiltonian simulation and phase estimation [43, 44]. Here, the idea is to apply phase estimation to the exponential \( \exp(-iH) \) with the prepared state \( \rho \) as input. Our goal is to estimate

\[
F(\rho, \rho_0) = \frac{1}{n} \sum_{j} q_j \left| \langle E_0|\rho_0 \rangle \right|^2 = \sum_{j} q_j F(\langle \psi_j|\rho_0 \rangle),
\]

where we have written \( \rho = \sum_{j} q_j |\psi_j \rangle \langle \psi_j| \) in its eigenbasis. Thus the fidelity of \( \rho \), a mixture of pure states, with \( |E_0 \rangle \) can be equivalently viewed as a mixture of fidelities of the component pure states \( |\psi_j \rangle \) with \( |E_0 \rangle \). Therefore it suffices to analyze phase estimation independently for each \( |\psi_j \rangle \). With each input state \( |\psi_j \rangle = \sum_{i=0}^{n} \psi_{ij} |E_i \rangle \) expressed in the eigenbasis of \( H \), the output of the phase estimation algorithm applied to \( \exp(-iH) \) is a state of the form \( |\phi_j \rangle = \sum_{i=0}^{n} p_{ij} |E_i \rangle \). Here, \( |E_i \rangle \) is a t-qubit state when measured in the computational basis yields the estimate \( \hat{E}_i \) of the eigenvalue \( E_i \) of \( H \) as its outcome. Performing a measurement with POVM elements \( \{|E_0 \rangle \langle E_0|, \text{id} - |E_0 \rangle \langle E_0|\} \) of the first register then allows one to estimate the desired fidelity \( F(\langle \psi_j|\rho_0 \rangle, |E_0 \rangle) \). Recombining the component fidelities according to their probability weights \( q_j \), yields the overall fidelity \( \sum_{j} q_j F(\langle \psi_j|\rho_0 \rangle, |E_0 \rangle) = F(\rho, |E_0 \rangle) \) as the observed frequency of the outcome \( E_0 \).

The accuracy of the estimate and the success probability are determined by \( t \). In order to distinguish the different possible measurement outcomes uniquely, we require that the estimate \( \hat{E}_i \) be accurate up to an additive error of \( 2^{-n} \ll \Delta \). For this to be the case with probability \( 1 - \beta \) we require the first register to comprise \( t = n + \log(2 + 1/\beta) \) qubits [44]. Again, using Hoeffding’s inequality, we find that in order to determine \( F(\rho, \rho_0) = \sum_{j} q_j |p_{j0}|^2 \) up to an additive error \( \epsilon \) with probability \( 1 - \alpha \), a number \( m \geq \ln(2/\alpha)/(2\epsilon^2) \) of i.i.d. measurements of \( |E_0 \rangle \langle E_0| \) on \( \rho \) are sufficient.

While such an approach would of course remove the precision bottleneck, requires few measurements and allows the certification of ground states of arbitrary Hamiltonians, it also puts significantly more requirements on capabilities of a prospective quantum simulator: it requires an additional \( n \)-qubit register and the capability to perform the phase estimation algorithm on the joint system. Essentially, a universal quantum computer is required for phase estimation. Ultimately, it will be up to the experimentalist to choose the most effective trade-off.

IV. SUMMARY AND DISCUSSION

In this work, we have discussed in what sense weak-membership variants of the certification of ground states of frustration-free Hamiltonians can be used to certify the correctness of analog quantum simulation and instances of quantum computation. For this certification to be possible, merely local Hamiltonian terms have to be measured, in a way that is perfectly experimentally possible on a number of platforms, in conjunction with efficient classical processing. No multi-round interactive proof systems is required for that. The results here challenge the view that is often expressed specifically in the physics literature, namely that in order to certify a quantum simulator, one needs to be able to efficiently keep track of the outcome of the simulation. Instead, in important cases, one can certify its functioning without being able to efficiently predict the actual result of the simulation. The discussion of universal quantum computing is in this context is primarily relevant to show that on a classical device computationally hard problems can indeed be solved by analog quantum simulators.

Having said this, the observations made here further motivate the quest of identifying further intermediate problems. Such an intermediate problem would satisfy two criteria:

1. The problem is believed to be intractable classically, but it is not necessarily the case that arbitrary quantum computations can be reduced to solving it.

2. A reliable quantum simulator is conceivable that can be practically realised with present day experimental capacities and solves that problem.

Identifying such problems requires a complexity-theoretic analysis of problems that are related to physical models. Notions of boson sampling offer interesting instances of intermediate problems [45–47]. As an illustration of our approach we have sketched how our method can be combined with the results of Ref. [11] to certify the preparation of a probability distribution, which is provably hard to simulate on any classical computer under a plausible complexity-theoretic assumption.

V. ACKNOWLEDGEMENTS

We would like to thank M. Gluza, D. Gosset, and B. M. Terry for discussions and the EU (AQuS, SIQS, RAQUEL), the COST network, the DFG (EI 519/7-1), the ERC (TAQ), the Studienstiftung des Deutschen Volkes, the Templeton Foundation, and the Humboldt Foundation for support. We also thank the anonymous referee to point out the more complex, but constant-precision alternative for fidelity estimation. Finally, we thank Roberto Di Candia for pointing us to a small error in Eqs. (2) and (5) in an earlier version of this work.
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