We studied the quantum adiabatic algorithm for combinatorial optimization. We obtained a simpler proof of the adiabatic theorem. However, we did not make much progress in developing new tools for rigorous analysis of the algorithm's performance on realistic optimization problems. This analysis seems to be substantially more difficult than for classical algorithms such as simulated annealing, because the ground state does not have a simple form.

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Report Title
QuaCGR Fellowship: Adiabatic Quantum Algorithms

ABSTRACT
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We also proved some interesting results about the complexity of the Local Consistency problem (deciding whether local density matrices that describe small pieces of a quantum system are consistent with a single overall state). In particular, we showed that this problem is QMA-complete. We also showed that N-representability, an important problem in quantum chemistry, is QMA-complete.

List of papers submitted or published that acknowledge ARO support during this reporting period. List the papers, including journal references, in the following categories:

(a) Papers published in peer-reviewed journals (N/A for none)

Number of Papers published in peer-reviewed journals: 1.00

(b) Papers published in non-peer-reviewed journals or in conference proceedings (N/A for none)
Number of Papers published in non peer-reviewed journals: 0.00

(c) Presentations

Number of Presentations: 1.00

Non Peer-Reviewed Conference Proceeding publications (other than abstracts):
Number of Non Peer-Reviewed Conference Proceeding publications (other than abstracts): 0

Peer-Reviewed Conference Proceeding publications (other than abstracts):


Number of Peer-Reviewed Conference Proceeding publications (other than abstracts): 2

(d) Manuscripts


Number of Manuscripts: 2.00
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### Student Metrics

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- The number of undergraduates funded by your agreement who graduated during this period and will continue to pursue a graduate or Ph.D. degree in science, mathematics, engineering, or technology fields: ...... 0.00
- Number of graduating undergraduates who achieved a 3.5 GPA to 4.0 (4.0 max scale): ...... 0.00
- Number of graduating undergraduates funded by a DoD funded Center of Excellence grant for Education, Research and Engineering: ...... 0.00
- The number of undergraduates funded by your agreement who graduated during this period and intend to work for the Department of Defense: ...... 0.00
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Number of Inventions:

Graduate Students PERCENT_SUPPORTED

Yi Kai Liu 1.00 FTE Equivalent: 1.00 Total Number: 1

Names of Post Doctorates

Names of Faculty Supported

Names of Under Graduate students supported

Student Metrics

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Names of Personnel receiving masters degrees

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### Sub Contractors (DD882)

### Inventions (DD882)
1 Problem Statement

We studied the quantum adiabatic algorithm for combinatorial optimization, first proposed by [8]. The algorithm works as follows. One constructs a time-dependent Hamiltonian \( H(t) \), \( 0 \leq t \leq T \), which can be efficiently implemented on a quantum computer. \( H(t) \) is designed so that, at time 0, the ground state of \( H(0) \) is known and can be prepared efficiently, while at time \( T \), the ground state of \( H(T) \) encodes the solution of some combinatorial optimization problem. Furthermore, \( H(t) \) changes “slowly” over time.

The idea is that, if we prepare the system in the ground state at time 0, and allow it to evolve until time \( T \), then the system will follow the ground state of \( H(t) \), and end up finding the solution to our optimization problem. This will work provided the change in \( H(t) \) is sufficiently slow; in particular, by the adiabatic theorem, it suffices to have \( T \gg \text{poly}(1/g_{\text{min}}) \), where \( g_{\text{min}} \) is the minimum gap between the two lowest eigenvalues of \( H(t) \) (for all times \( t \)). Thus, the spectral gap of \( H(t) \) essentially determines the running time of the algorithm. (Other factors, such as the norm of \( H(t) \) and its derivatives, also play a role, but they are usually less significant.)

For example, suppose we want to minimize the function \( f \) on the Boolean hypercube \( \{0, 1\}^n \). The adiabatic algorithm might use a Hamiltonian of the form:

\[
H(t) = (1 - t/T)H_0 + (t/T)H_1
\]

\[
H_0 = \sum_{i=1}^{n} \sigma_x^{(i)}
\]

\[
H_1 = \sum_{z_1, \ldots, z_n \in \{0, 1\}} f(z_1, \ldots, z_n) |z_1\rangle\langle z_1| \otimes \cdots \otimes |z_n\rangle\langle z_n|.
\]

Here, the system consists of \( n \) qubits, \( \sigma_x^{(i)} \) denotes the \( \sigma_x \) Pauli operator acting on the \( i \)’th qubit, and \( |0\rangle \) and \( |1\rangle \) denote the up and down \( \sigma_z \) eigenstates for a single qubit.

*Computer Science and Engineering, University of California, San Diego. E-mail: y9liu@cs.ucsd.edu, yikai@caltech.edu.
The adiabatic algorithm was originally proposed to solve problems such as 3-SAT, but no rigorous analysis of the running time is known in that case. Rigorous analyses are only available for very simple “toy problems” (see, e.g., [22, 7, 18]). These give some insight into the algorithm—in particular, they show how the adiabatic algorithm can succeed by tunnelling through potential barriers, and how it can fail by getting drawn into local minima. But it remains difficult to understand the performance of the adiabatic algorithm on real optimization problems, because the situation there is so much more complicated. At best, there are heuristic arguments for those cases (see, e.g., [20]).

Our goal was to develop new techniques for rigorous analysis of the adiabatic algorithm, with the hope of understanding the performance of the algorithm on some moderately complicated, quasi-realistic problems. A rigorous analysis of this type would be useful in confirming (or refuting) the conclusions obtained from heuristic arguments. We remark that this sort of analysis has been carried out, to some extent, for classical algorithms such as simulated annealing (see, e.g., [21, 9, 1, 5, 6, 4]).

2 Summary of Results

The following is a summary of our main results. For more details, see the annual progress reports, and the reprints of papers that were submitted to ARO.

2.1 The Adiabatic Algorithm

We obtained a simple proof of the adiabatic theorem, using a martingale argument [11]. Another more powerful proof was discovered independently by Ambainis and Regev [2]. These results give some intuition about why the spectral gap is significant: a small spectral gap corresponds to a rapid change in the ground state of the Hamiltonian, which in turn implies a longer running time, to ensure that the system remains in the ground state. In many cases, the system seems to undergo a quantum phase transition, where there is an “avoided crossing” between the two lowest energy eigenvalues, coinciding with a sudden change in the ground state.

Unfortunately, we made little progress in developing new tools for rigorous analysis of the problem. The main difficulty is that, generally speaking, the ground state of the Hamiltonian does not have a simple, explicit description. The only known exceptions are when: (1) the Hamiltonian has a symmetry that effectively reduces the dimension of the problem, e.g., it is symmetric under permutations of the qubits [8, 22]; (2) the Hamiltonian has a special structure that can be diagonalized exactly, e.g., it can be transformed into a system of noninteracting fermions [18]; or (3) the ground state has relatively uncomplicated correlations and entanglement, e.g., certain 1-D spin chains that can be analyzed using the martingale method [17].

For comparison, note that in classical simulated annealing, the stationary distribution of
the Metropolis random walk does have a simple description, even in a complicated system—it is the Gibbs distribution. This makes it possible to calculate the second eigenvalue, for instance by variational methods, and thus bound the spectral gap.

Perhaps heuristic arguments and numerical methods, developed in condensed matter physics, are a more viable approach to understanding the adiabatic algorithm. For instance, Schutzhold and Schaller have an interesting interpretation of the algorithm in terms of first- and second-order quantum phase transitions [19]. Also, it may be possible to simulate some instances of the adiabatic algorithm using non-exact methods such as matrix-product states [3].

2.2 The Local Consistency Problem

We also obtained several results on the complexity of the “Local Consistency” problem. (This is not related to the adiabatic algorithm, but it is nonetheless interesting.) The problem is as follows: given a collection of density matrices \( \rho_1, \ldots, \rho_m \), where \( \rho_i \) describes a subset of qubits \( C_i \) in an \( n \)-qubit system, with \( |C_i| \leq O(1) \), decide whether there exists an \( n \)-qubit state \( \sigma \) that agrees with all of the \( \rho_i \) on the subsets \( C_i \).

In a certain sense, this problem is complementary to the Local Hamiltonian problem (finding the ground state energy of a local Hamiltonian), which is QMA-complete [10]. We showed that Local Consistency is QMA-complete, via a novel reduction using convex optimization with a membership oracle [12]. We also showed that \( N \)-representability, an important problem in quantum chemistry, is QMA-complete [16]. In addition, we proved a structural property of Local Consistency (that if a solution exists, it can be chosen to be a Gibbs state) [13]; and we identified special cases of Local Consistency, for 1-D and stoquastic systems, whose complexity appears to be strictly easier than QMA [15]. These results are described in greater detail in my thesis [14].

2.3 Publications and Demographic Data

The following information is for the period from 8/1/2006 to 8/31/2007 (since last year’s progress report). Papers during this reporting period: [16] (published in journal), [14] (PhD dissertation). Demographic data: 1 full-time graduate student supported by this agreement. Inventions: none. Technology transfer: none. Other activities: Yi-Kai Liu gave a talk at RANDOM 2006; visited the Institute for Quantum Computation (IQC) at Waterloo in January 2007; gave an invited talk at QIP 2007; and attended the SQuInT 2007 workshop.

References


The central theoretical problem in the field of many-body strongly correlated quantum systems is to find efficient ways of simulating Schrödinger’s equations: it is very easy to write down those equations, but notoriously difficult to solve them or even to find approximate solutions. The main difficulty is the fact that the dimension of the Hilbert space describing a system of $N$ quantum particles scales exponentially in $N$. This makes a direct numerical simulation intractable: every time an extra particle is added to the system, the computational resources would have to be doubled.

The situation is not hopeless, however, as in principle it could be that all physical wavefunctions, i.e., the ones that are realized in nature, have very special properties and can be parameterized in an efficient way. The idea would then be to propose a variational class of wavefunctions that capture the physics of the systems of interest, and then do an optimization over this restricted class. This approach has proven to be very successful, as witnessed by mean field theory and renormalization group methods. So far, an efficient variational class to describe complex wavefunctions such as those arising in quantum chemistry has not been found.

One of the basic problems in quantum chemistry is to find the ground state of a Hamiltonian describing the many-body system of an atom or molecule. The essential element that makes typical Hamiltonians very ungeneric is the fact that at most 2-body interactions occur. This implies that the number of free parameters in such Hamiltonians scales at most quadratically in the number of particles or modes, and hence the ground states of all such systems form a small-dimensional manifold.

In the case of a Hamiltonian with only 2-body interactions, the energy corresponding to a wavefunction is computed by a wavefunction that is consistent with some global state over $N$ fermions. This makes a direct numerical simulation intractable: every time an extra particle is added to the system, the computational resources would have to be doubled.

The dual problem of determining the ground state energy of a local Hamiltonian is to decide whether all physical wavefunctions, i.e., the ones that are realized in nature, have very special properties and can be parameterized in an efficient way. The idea would then be to propose a variational class of wavefunctions that capture the physics of the systems of interest, and then do an optimization over this restricted class. This approach has proven to be very successful, as witnessed by mean field theory and renormalization group methods. So far, an efficient variational class to describe complex wavefunctions such as those arising in quantum chemistry has not been found.

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a given set of local density operators can be realized as the reduced density operators of the same global state. Checking the consistency of such a set of local density operators \( \rho^{(j)} \), i.e., checking whether there exists a global state \( \sigma \) compatible with those local density operators, is a QMA-complete problem itself [14]. In the present paper, we will use very similar techniques to prove that \( N \)-representability, which is the fermionic version of that problem, is also QMA-complete.

To sketch the main ideas of the proof, we will first consider the classical marginal problem: suppose we have \( N \) random variables that can take the values \( \pm 1 \), according to some joint probability distribution, and we define the 2-variable marginal distributions

\[
p(a_i, a_j) = \sum_{a_1, a_2, \ldots, a_N} p(a_1, a_2, \ldots, a_N).
\]

Given a set of marginal probability distributions, we would like to check whether they are consistent, i.e., whether there exists a global probability distribution whose marginals are equal to the ones we were given.

Suppose that there exists an efficient algorithm (whose running time is polynomial in \( N \)) to solve this problem. Then it would be possible to identify ground states of all Ising spin glasses. The strategy would be as follows: since the set of consistent marginals is convex and the energy is a linear function of the marginals, the problem amounts to minimizing a linear function subject to a set of convex constraints that can be checked in polynomial time. This can be done in polynomial time using the ellipsoid method [11]. It therefore follows that an efficient algorithm for checking consistency allows to find ground states of Ising spin glasses, a problem that is known to be NP-hard [12]. Hence the problem of determining whether a set of binary marginals is consistent is itself NP-hard.

Let us now formulate the \( N \)-representability problem in the context of quantum chemistry. Electrons and nuclei tend to arrange themselves such as to minimize their energy, and the binding energy of a molecule can be determined by calculating the minimal energy of the corresponding Hamiltonian. In practice, the nuclei are fairly well localized and can be treated as classical degrees of freedom, and the wavefunction of the \( N \) electrons can be approximated as a linear combination of tensor products of the \( d \) single-particle modes in the system (those form the basis set).

As electrons are fermionic, the complete wavefunction must be antisymmetric, and this is most easily taken into account by working in the formalism of second quantization:

\[
|\psi\rangle = \sum_{i_1, \ldots, i_d = 0}^{1} c_{i_1, \ldots, i_d} (a_j^\dagger)^{i_1} \cdots (a_j^\dagger)^{i_d} |\Omega\rangle.
\]

Here \( a_j^\dagger \) is the creation operator for the \( j \)’th mode, and \( |\Omega\rangle \) represents the vacuum state without fermions. The creation and annihilation operators obey the following anticommutation relations:

\[
\{a_i, a_j\} = 0 = \{a_i^\dagger, a_j^\dagger\} \quad \{a_i, a_j^\dagger\} = 2\delta_{ij}.
\]

Note that we restrict ourselves to the subspace of states with exactly \( N \) fermions. \( d \) denotes the number of modes, which is typically much larger than \( N \). The number of degrees of freedom is \( \binom{d}{N} \), which grows exponentially in \( N \) when \( d \geq cN \) for some constant \( c > 1 \).

In the case of quantum chemistry, the Hamiltonian typically contains only one- and two-body interactions between all modes, and so it can be written as a linear combination of terms of the form \( a_i^\dagger a_j \) and \( a_i^\dagger a_j^\dagger a_k a_l \).

The \( 2 \)-fermion reduced density matrix (2-RDM) is calculated by tracing out all but two of the fermions:

\[
\rho^{(2)} = \text{tr}_{N-2} \rho^{(N)}.
\]

where \( \rho^{(N)} \) is a mixture of states \( |\psi\rangle \) with exactly \( N \) fermions. In the language of second quantization, the matrix elements of the 2-RDM are given by:

\[
\rho^{(2)}_{ijkl} = \frac{1}{N(N-1)(a_i^\dagger a_j^\dagger a_k a_l)}.
\]

The \( N \)-representability problem (with \( d \) modes) can now be stated as follows. Consider a system of \( N \) fermions and \( d \) modes, \( d \leq \text{poly}(N) \). (For purposes of complexity, we consider \( N \) to be the “size” of the problem.) We are given a 2-fermion density matrix \( \rho \), of size \( \frac{d(d-1)}{2} \times \frac{d(d-1)}{2} \), where each entry is specified with \( \text{poly}(N) \) bits of precision. In addition, we are given a real number \( \beta \geq 1/\text{poly}(N) \), specified with \( \text{poly}(N) \) bits of precision. The problem is to distinguish between the following two cases:

- There exists an \( N \)-fermion state \( \sigma \) such that \( \text{tr}_{N-2} \rho^{(2)}(\sigma) = \rho \). In this case, answer “YES.”
- For all \( N \)-fermion states \( \sigma \), \( \|\text{tr}_{N-2} \rho^{(2)}(\sigma) - \rho\|_1 \geq \beta \).

In this case, answer “NO.”

If neither of these cases applies, then one may answer either “YES” or “NO.” Note that we do not insist on solving the problem exactly; we allow an error tolerance of \( \beta \geq 1/\text{poly}(N) \). We use the \( \ell_1 \) matrix norm or trace distance, \( \|A\|_1 = \text{tr} |A| \), to measure the distance between \( \sigma \) and \( \rho \).

We will show that \( N \)-representability is QMA-complete. The proof consists of two parts. First, we show that any 2-local Hamiltonian of spins can be simulated using a 2-local Hamiltonian of fermions with \( d = 2N \). Using techniques of convex programming, we show that an oracle for \( N \)-representability would allow us to estimate the ground state energies of 2-local Hamiltonians; thus, \( N \)-representability is QMA-hard.

Second, we show that \( N \)-representability is in QMA; specifically, we construct a quantum verifier that can
check whether a 2-particle state is \( N \)-representable, given a suitable witness.

Let us first show how to map a 2-local Hamiltonian, defined on a system of \( N \) qubits, to a 2-local Hamiltonian on fermions, with \( d = 2N \) modes; this is the opposite of what has been done in \cite{13}. The idea is to represent each qubit \( i \) as a single fermion that can be in two different modes \( a_i, b_i \); so each \( N \)-qubit basis state corresponds to the following \( N \)-fermion state:

\[
| z_1 \rangle \otimes \cdots \otimes | z_N \rangle \mapsto (a_1^\dagger)^{1-z_1}(b_1^\dagger)^{z_1} \cdots (a_N^\dagger)^{1-z_N}(b_N^\dagger)^{z_N}| \Omega \rangle.
\]

(1)

Also, all the relevant single-qubit operators should correspond to bilinear functions of the creation and annihilation operators (this construction guarantees that operators on different qubits commute). This lets us simplify the fermionic Hamiltonian as follows. Since we are only interested in how it acts on \( N \)-particle states, we can write \( a_i^\dagger a_j = \frac{1}{\sqrt{2}} \alpha_j^\dagger (\sum_k a_k^\dagger a_k)a_j \). So we can assume that all the terms in the Hamiltonian are of the form \( a_j^\dagger a_j a_k \).

Second, convex optimization algorithms usually require that the set \( K \) of feasible solutions be full-dimensional, i.e., \( K \) cannot lie in a lower-dimensional subspace. So we have to represent the state \( \rho \) in such a way that there are no redundant variables.

We write our Hamiltonian in the form \( H = \sum_{S \in S} \gamma_S S \) (plus a constant term); the coefficients \( \gamma_S \) can be computed using the Gram-Schmidt procedure. It is easy to see that \( \text{tr}(H \rho) = \sum_{S \in S} \gamma_S \alpha_S \). Our convex program is as follows: find some \( \tilde{\alpha} \in K \) that minimizes \( f(\tilde{\alpha}) = \sum_{S \in S} \gamma_S \alpha_S \).

We can solve this convex program in polynomial time, using the shallow-cut ellipsoid algorithm \cite{11}, and references cited therein). We mention a few technical details. The algorithm requires some additional information about \( K \), namely a guarantee that \( K \) is contained in a ball of radius \( R \) centered at 0, and \( K \) contains a ball of radius \( r \) centered at some point \( p \). (Also, the running time of the algorithm grows polynomially in \( \log(R/r) \).

In our case, we can set \( R = \sqrt{\ell} \), and \( r = 1/\text{poly}(\ell) \).
solution to the convex program, in order to solve the Local Hamiltonian problem. It turns out that $1/poly(N)$ precision is sufficient. The ellipsoid algorithm still works in this setting; see \cite{43} for details.

Note that, in place of the ellipsoid algorithm, we could have used a different algorithm based on random walks in convex bodies \cite{44}; this was the approach used in \cite{45}. However, it is not clear if we could use one of the interior-point methods for convex optimization; these methods are substantially faster, but they usually require an explicit description of the constraints, not just a membership oracle.

This completes the proof that $N$-representability is QMA-hard. As a corollary, we have also proven that estimating the ground state energy for local fermionic Hamiltonians is QMA-hard. Note that the use of the ellipsoid algorithm to reduce a convex optimization problem to a convex membership problem is not new; see \cite{44} for references to related work.

Next, we show that $N$-representability is in QMA. That is, we construct a poly-time quantum verifier $V$ that takes two inputs: a description of the problem (that is, $\rho$ and $\beta$); and a “witness” $\tau$, which is a quantum state on polynomially many qubits. The verifier $V$ should have the following property: if $\rho$ is $N$-representable, there exists a witness $\tau$ that causes $V$ to output “true” with probability $\geq p_1$; if $\rho$ is not $N$-representable (within error tolerance $\beta$), then for all possible states $\tau$, $V$ outputs “true” with probability $\leq p_0$; and $p_1 - p_0 \geq 1/poly(N)$.

The idea is that, when $\rho$ is $N$-representable, the correct witness $\tau$ consists of (multiple copies of) an $N$-fermion state $\sigma$ that satisfies $tr_{3,...,N}(\sigma) = \rho$. Then the verifier can use quantum state tomography to compare $\sigma$ and $\rho$.

We represent the $N$-fermion state $\sigma$ using $d$ qubits, via the following mapping:

$$(a_1^\dagger)^{i_1}...\cdot(a_d^\dagger)^{i_d}\cdot|0\rangle \mapsto |i_1\rangle \otimes ... \otimes |i_d\rangle.$$ Call the resulting qubit state $\hat{\sigma}$. We use the Jordan-Wigner transform to map the fermionic annihilation operators to qubit operators:

$$a_i \mapsto A_i = - (\otimes_{k<i} \sigma_k^x) \otimes |0\rangle \langle 1|.$$ Thus, an observable $O = a_j^\dagger a_k^\dagger a_k a_j$ is transformed into $\hat{O} = A_j^\dagger A_k^\dagger A_k A_j^\dagger + A_j^\dagger A_k^\dagger A_j A_k$, which is a tensor product of many single-qubit observables and one four-qubit observable.

We claim that the expectation value $\langle \hat{O} \rangle$ can be estimated efficiently. Without loss of generality, assume that the eigenvalues of $\hat{O}$ lie in the interval $[0,1]$. Then there is an efficiently-implementable measurement which outputs “1” with probability $\langle \hat{O} \rangle$ \cite{45}. By repeating this measurement on multiple copies of the same state, we can estimate $\langle \hat{O} \rangle$; for our purposes, it is enough to have polynomially many copies of the state.

We now describe the verifier $V$. The witness $\tau$ consists of several (i.e., polynomially many) blocks, where each block has $d$ qubits, supposedly representing one copy of the state $\hat{\sigma}$. On each block, $V$ measures the observable $\sum_k |1\rangle \langle 1|_k$, and if the outcome does not equal $N$, $V$ outputs “false.” This projects each block onto the space of $N$-particle states.

Next, $V$ performs measurements on each block, to estimate the expectation values of $\hat{\sigma}$, for a suitable set of observables. Then $V$ checks whether they match the expectation values of $\rho$. One problem arises: the prover could try to cheat by entangling the different blocks of qubits. One can show that this does not fool the verifier, using a Markov argument, as was done in \cite{45}. This suffices to show that $N$-representability is in QMA, and hence finishes the proof.

What can be said about the complexity of the pure-state $N$-representability problem, where one has to decide whether the reduced density operators arise from a pure state of $N$ fermions? In that case, the verifier must be able to convince himself that the state he gets is pure. This can be done when he gets two states $\rho$ and $\sigma$ that are promised to be uncorrelated, i.e. that he gets the state $\rho \otimes \sigma$: then Arthur can calculate the expectation value of the observable $tr(\rho \sigma)$, and this can only be close to 1 when $\rho$ is pure and $\sigma \approx \rho$. Indeed, if $tr(\sigma^2) \leq 1 - \epsilon$, then for all states $\tau$, $tr(\tau \sigma) \leq \sqrt{tr(\sigma^2)tr(\tau^2)} \leq (1-\epsilon)^{1/2} \leq 1 - \epsilon/2$. The problem however is that Merlin can cheat, and hand out a correlated state to Arthur, such that the above test becomes inconclusive. This is precisely the feature that distinguishes the complexity class QMA(k) from QMA \cite{10}: in QMA(k), the verifier is promised to get a tensor product of different states, and it has been conjectured that QMA(k) is strictly larger than QMA. The above discussion shows that the pure-state $N$-representability problem is contained in QMA(k), and it is hence plausible that it is harder than the $N$-representability problem. It would be very interesting to investigate whether the problem is QMA(k)-complete.

It is remarkable that checking consistency of 2-body reduced density operators of fermionic states is so hard, while checking consistency of 1-body reduced density operators is simple \cite{2}. This can be easily understood from the previous discussion: the extreme points of the convex set of 1-body density operators $\langle a_i^\dagger a_j \rangle$ correspond to ground states of Hamiltonians only containing bilinear terms in $a_i^\dagger$ and $a_j$; such Hamiltonians can easily be diagonalized as they represent systems of free fermions, and hence the consistency problem can easily be solved. As shown in \cite{2}, consistency can be decided in that case based solely on the eigenvalues of the reduced density operators. A number of related problems, where consistency only depends on the eigenvalues, have been investigated recently \cite{17}; most remarkably, a characteri-
sation has been obtained for the polytope of 1-particle marginals that are $N$-representable under the condition that the $N$-particle wavefunction must be pure.\cite{18}

These results have to be contrasted with our problem of finding the $N$-representable 2-body density operators, where the eigenvalues alone are not enough to decide consistency but also the eigenvectors are relevant. Actually, let us consider the simpler problem of deciding $N$-representability of 2-fermion density operators where only the diagonal elements $D_{ij} = \langle a_i^\dagger a_j^\dagger a_j a_i \rangle$ are specified. If we consider the case $d = 2N$ and the mapping discussed above, one easily finds that the extreme points of this set would be obtained by ground states of local spin Hamiltonians which only contain commuting $\sigma^z$ operators. These correspond to spin-glasses, and so the problem of deciding $N$-representability of $\{D_{ij}\}$ is NP-hard.\cite{12}. It was indeed pointed out a long time ago that $N$-representability restricted to the diagonal elements is equivalent to a combinatorial problem\cite{11} that was later shown to be equivalent to the NP-hard problem of deciding membership in the boolean quadric polytope.\cite{20}.

Let’s finally discuss the relevance of the above results in the context of quantum chemistry. We have shown that it is a hopeless task to determine the ground states of all local fermionic Hamiltonians, and in particular, that an approach by means of the $N$-representability problem is intractable, even on a quantum computer. But it is possible that other physical systems, e.g., systems with different particle statistics, additional symmetry or a limited number of modes, might allow for an efficient characterisation of the two-particle reduced density matrices, and hence an efficient calculation of the ground states of local Hamiltonians. This seems to be the case in e.g. one-dimensional translational invariant spin systems, where the density matrix renormalization group\cite{21} allows for a systematic approximation of the allowed convex set of reduced density operators from within\cite{22}. A very different numerical approach has been developed in the context of quantum chemistry and it is known as the contracted Schrödinger equation.\cite{4}. In that method, one approximates the convex set from the outside, and the $N$-representability problem is a crucial ingredient of the algorithm. The foregoing discussion shows that this approach has to break down in the most general case, and it would be very interesting to investigate the conditions under which those approximations are justified.

In conclusion, we investigated the problem of $N$-representability, and characterized its computational complexity by showing that it is QMA-complete. Obviously, the theory of quantum computing was a prerequisite to pinpoint the computational complexity of this classic problem.

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\bibitem{21} U. Scholz, Rev. Mod. Phys. 77, 259 (2005).
\bibitem{23} Technically, the problem must be formulated as a yes-no question, e.g., “for a given Hamiltonian $H$, is the ground state energy less than $\gamma$?” If the answer is yes, there exists a “witness” (e.g., the ground state) which can prove it. Moreover, there is an efficient algorithm that can check the validity of a witness.
\end{thebibliography}
Another option would be to represent each qubit $i$ using either zero or two fermions, e.g., for a single qubit, $|0\rangle$ and $|1\rangle$ correspond to $|\Omega\rangle$ and $a_i^\dagger b_i^\dagger |\Omega\rangle$. Define the mapping $\sigma_i^\tau \leftrightarrow \begin{pmatrix} a_i^\dagger b_i + b_i a_i^\dagger \\ b_i a_i + a_i b_i^\dagger \end{pmatrix}$. Let $\sigma_i^\tau \leftrightarrow \begin{pmatrix} a_i b_i^\dagger - a_i^\dagger b_i \\ a_i^\dagger b_i - b_i a_i \end{pmatrix}$. Then the Hamiltonian would act identically on the subspace with one fermion per site, and on the subspace with zero or two fermions per site, and even a small constant in front of the projectors $P_i$ would be enough to guarantee that we end up in the right subspace.

One possible set of observables is as follows: First, define $a_i = a_i a_i^\dagger$, for all pairs $I = \{i_1, i_2\}$, $i_1 < i_2$. Also fix an ordering on the pairs $I$. We now define the following observables: $X_{IJ} = a_i^\dagger a_j + a_j^\dagger a_i$, for all $I < J$; $Y_{IJ} = -ia_i^\dagger a_j + ia_j^\dagger a_i$, for all $I < J$; and $Z_I = a_i^\dagger a_i$, for all $I$. These operators are Hermitian, with eigenvalues in the interval $[-1, 1]$. Taking real linear combinations, these observables form a basis for the space of 2-fermion density matrices.

It is easy to see that $K$ is contained in a ball of radius $R = \sqrt{t}$, since for all $\vec{a}$, we have $-1 \leq \langle \vec{a} | \sigma | \vec{a}\rangle \leq 1$. The second claim, that $K$ contains a ball of radius $r = 1/\text{poly}(t)$, is less trivial. The proof is as follows.

We will consider $N$-representability for different values of $N$; let $K_N$ denote the set of all vectors $\vec{a}$ that are $N$-representable. Obviously, $K_2$ contains a ball of radius $1/\text{poly}(t)$ (this is the trivial case). We will prove that $K_N$ contains a ball of radius $1/\text{poly}(t)$, for all $2 \leq N \leq d - 2$.

We define “particle-hole” observables, by replacing $a_i$ with $a_i^\dagger$, and versa: $X'_{IJ} = a_i^\dagger a_j^\dagger + a_j^\dagger a_i^\dagger$; $Y'_{IJ} = -ia_i^\dagger a_j^\dagger + ia_j^\dagger a_i^\dagger$; and $Z'_I = a_i^\dagger a_i$. Let $\vec{a}'$ denote a vector containing expectation values for these observables; let $K_N'$ be the set of all $\vec{a}'$ that are $N$-representable.

First, we claim that $K_2 = K_2'$. Notice that there is a natural correspondence between the 2-particle Slater basis states and the $(d-2)$-particle Slater basis states: the 2-particle state with modes $i$ and $j$ occupied corresponds to the $(d-2)$-particle state with modes $i$ and $j$ empty. So take any point $\alpha \in K_2$, which represents the expectation values of the 2-particle observables for some 2-particle state $\sigma$. Use $\sigma$ to construct a $(d-2)$-particle state $\tau$, by replacing each 2-particle Slater basis state with the corresponding $(d-2)$-particle Slater basis state. Then the expectation values of the 2-particle observables for $\sigma$ are exactly the expectation values of the 2-hole observables for $\tau$. So $\alpha$ is in $K_2'$. This shows that $K_2 \subseteq K_2'$. A similar argument shows that $K_2' \subseteq K_2$. So $K_2 = K_2'$.

Next, we claim that there is an invertible linear transformation $A$ that maps $K_{d-2}$ to $K_{d-2}$. First we map $K_{d-2}'$ to $K_{d-2}$. Observe that we can write each 2-particle operator as a linear combination of 2-hole operators. (This holds provided we restrict the operators to act only on the subspace of $(d-2)$-particle states.) For instance, when $I \cap J = \emptyset$, $a_i^\dagger a_j = a_j^\dagger a_i$. When $|I \cap J| = 1$, we write equations such as $a^\dagger_i a^\dagger_j a_i a_j = a^\dagger_j a^\dagger_i a_j a_i = -a^\dagger_i a^\dagger_j a_j a_i = a^\dagger_j a^\dagger_i a_i a_j$; then use the fact that $a_i a_j = \sum_{\ell \neq i} a_i a^\dagger a^\dagger a_i a_j a_i = \sum_{\ell \neq j} a^\dagger a^\dagger a_i a^\dagger a^\dagger a_j a_i$. When $I = J$, we have $a_i^\dagger a_i = \sum_{\ell \neq i} a^\dagger a^\dagger a_i a^\dagger a^\dagger a_i a_i$. (For each of these equations, it is easy to check that the left and right sides act identically on all $(d-2)$-particle Slater basis states.)

Thus, for any $(d-2)$-particle state $\sigma$, the expectation values of the 2-particle observables are linear functions of the expectation values of the 2-hole observables. Thus we have a linear transformation $A$ that maps $K_{d-2}'$ to $K_{d-2}$.

Similarly, we can map $K_{d-2}$ to $K_{d-2}$. We write each 2-hole operator as a linear combination of 2-particle operators (again, restricting the operators to act only on the subspace of $(d-2)$-particle states). For instance, when $I \cap J = \emptyset$, $a_i^\dagger a^\dagger_j a_i a^\dagger_j$. When $|I \cap J| = 1$, we write equations such as $a_i^\dagger a^\dagger_j a_i a^\dagger_j = a^\dagger_j a^\dagger_i a_i a^\dagger_j = a^\dagger_j a^\dagger_i a_i a^\dagger_j$. When $I = J$, we have $a_i^\dagger a_i = \sum_{\ell \neq i} a^\dagger a^\dagger a^\dagger a^\dagger a_i a_i$. (For each of these equations, it is easy to check that the left and right sides act identically on all $(d-2)$-particle Slater basis states.)