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Computational complexity in electronic structure

James Daniel Whitfield,*abc Peter John Loved and Alán Aspuru-Guzikut

In quantum chemistry, the price paid by all known efficient model chemistries is either the truncation of the Hilbert space or uncontrolled approximations. Theoretical computer science suggests that these restrictions are not mere shortcomings of the algorithm designers and programmers but could stem from the inherent difficulty of simulating quantum systems. Extensions of computer science and information processing exploiting quantum mechanics has led to new ways of understanding the ultimate limitations of computational power. Interestingly, this perspective helps us understand widely used model chemistries in a new light. In this article, the fundamentals of computational complexity will be reviewed and motivated from the vantage point of chemistry. Then recent results from the computational complexity literature regarding common model chemistries including Hartree–Fock and density functional theory are discussed.

1 Introduction

Quantum chemistry is often concerned with solving the Schrödinger equation for chemically-relevant systems such as atoms, molecules, or nanoparticles. By solving a differential and/or eigenvalue equation, the properties of the system and the dynamics of the state are obtained. Examples of properties include equilibrium geometries, the dissociation energy of molecules, and the vibrational frequencies.

The difficulty stems from the accuracy required and the apparent exponential growth of the computational cost with both the number of electrons and the quality of the description of the system. For
practical applications, the accuracy required is typically orders of magnitude smaller than the total energy of the system. As a concrete example, the Carbon atom has total electronic energy of about 37.8 Hartrees while the energy of a Carbon–Hydrogen bond is only 0.16 Hartrees. Solving the full eigenvalue equation takes on the order of \( n^3 \) operations for an \( n \times n \) matrix. However, when describing interacting many-electron systems, the dimension of the matrix increases exponentially with the number of electrons.

Consequently, the computational methods of electronic structure in chemistry are aimed at circumventing exact diagonalization, in this context called the full configuration interaction method. Avoiding exact diagonalization has led to a wide range of computational methods\(^1\)\(^–\)\(^4\) for computing properties of chemical interest. These methods have recently\(^5\)\(^–\)\(^9\) begun to include quantum simulation following Feynman’s suggestion\(^10\) to use quantum computers as simulators. Subsequent development of these ideas in quantum chemistry has led to new proposed methods utilizing quantum computational techniques\(^11\)\(^–\)\(^14\) and proof-of-principle experiments.\(^15\),\(^16\) However, questions about when and where one would expect a quantum computer to be useful\(^17\),\(^18\) have not been fully answered. Ideally, computational complexity can provide some answers about when, where, and why quantum computers would be useful for chemistry. At the same time, it could also help formalize intuitive understanding of when we can expect reliable results from traditional computational methods.

Many results in computational complexity have interchanged classical and quantum computers fluidly leading to new results on the complexity of computing properties of quantum systems.\(^19\) We review some recent results appearing in the computational complexity literature that touch on why electronic structure calculations are difficult. Our hope is to encourage future investigations into quantitative understanding of difficult instances of electronic structure calculations.

A similar, but much shorter, discussion of computational complexity in quantum chemistry by Rassolov and Garashchuk\(^10\) appeared in 2008 and provided many conjectures that have since been proven or extended.

This perspective assumes exposure to second quantization and mixed states in quantum mechanics. Standard bra-ket notation is used when referring to quantum states. All the necessary concepts from computational complexity and quantum computation are briefly introduced and motivated to make the article as self-contained as possible. A key omission from the present review is a number of computational complexity results\(^21\),\(^22\) concerning quantum-information based wave function ansatzes such matrix product states, density matrix renormalization group and their generalizations.\(^23\) These methods are becoming accepted into the mainstream of computational chemistry,\(^24\)\(^–\)\(^26\) but do not yet have the widespread availability of the selected methods included in the present work.

1.1 Overview and goals of this perspective article

The objective of this perspective is to provide the interested theoretical chemists access to the emerging subfield of computational complexity focused on classifying quantum Hamiltonians and approximants. We wish to provide new insights into quantum chemistry by encouraging an examination of the essence of computational model chemistries and its relation to the numerical simulations performed routinely in physical chemistry. To this end, we survey recent results in computational complexity that concern the methods used in the computational chemistry community. Admittedly, such a task is ambitious, but our hope is to provide a useful entry point to the growing literature by presenting a digestible account of key results to-date.

Following this overview, the article continues with an introduction to computer science for chemists. We try to use examples from physical chemistry to make the key ideas of theoretical computer science concrete and motivate the technical results presented later. We also provide a discussion of accuracy which is essential for interpreting the complexity of numerical or experimental results in a useful way.

Section 3 focuses on links between quantum chemistry and quantum physics which will be useful as the article progresses. We recall relevant aspects of quantum chemistry and draw connections to spin systems. This is essential as many of the early results in quantum computational complexity sprang out of investigations of spin systems which were likely candidates for constructing a quantum computer. Those results have since been extended in many directions, but in this article we focus on the difficulty of computing the ground state energy of quantum Hamiltonians.

In Section 4, our discussion of computation complexity in quantum chemistry begins, in earnest, with a discussion of Hartree–Fock and related mean-field approximations. The runtime of Hartree–Fock can be much worst than the heuristic arguments about the computation of the four-center integrals would suggest. This behavior is exemplified by calculations involving transition metals or actinide species where convergence of self-consistent solutions can be difficult to achieve. The crux of the argument is that Hartree–Fock could be used to solve for the ground state of certain spin systems which are known to belong to a difficult computational class of problems.

Next, we focus on approaches that do not involve the wave function as the fundamental variable. The first of these two

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approaches, discussed in Section 5, uses the two-body reduced density matrix. With the correct reduced density matrix, the evaluation of the ground state energy is straightforward. However, computing the criteria for valid density matrices is, in the worst cases, computational intractable. Here, the difficulty is demonstrated by showing, if the criterion were simple, then computations that are identified as difficult could be completed easily.

The final method of quantum chemistry to be expounded upon is density functional theory in Section 6. This is also the most nuanced result presented, in that its relevance to traditional quantum chemistry is weakened by relying on local magnetic fields. Using only the one-electron probability density as a computational variable requires that one introduces an unknown universal functional in order to obtain a practical route to computing the ground state energy. These unknown functionals are shown to belong to the same class of computational problems as computing the validity criteria for reduced density matrices. This is done, once more, by mapping a problem that is known to be computational intractable to the problem of computing the value of the universal functional. However, in this case, there is a series of two intermediate problems whose complexity is used to show that density functional theory is computationally difficult.

In Section 7, we provide a cursory discussion of other standard methods in quantum chemistry which were not covered earlier. Either we will relate them to the techniques discussed in this article or we point to what is known so far. Specifically, we briefly discuss multi-configuration wave function approaches, perturbation techniques, quantum Monte Carlo, de Broglie–Bohm, and time-dependent density functional theory.

In the penultimate section, we provide further technical remarks about error tolerance that are aimed at providing an introduction to contemporary computational complexity questions about extending what is known about the hardness of approximation to the quantum regime.

By the reader reaches the final section, we hope that he or she appreciates the extension of the theoretical foundations of quantum chemistry offered by computational complexity. To increase the usability of the article for a typical reader of PCCP, we give an outlook for computational chemists in Section 9. We hope that this article will motivate further investigation of the connections between chemical physics and quantum computational complexity theory. To facilitate future inquiry, we underline several important open questions.

2 Worst-case computational complexity for chemists

The purpose of this section is to provide a quick but precise introduction to computational complexity concepts. The aim is to set the stage for the results subsequently reviewed.

Computational complexity is the study of how resources required to solve a problem change with its size. For instance, space complexity is the scaling of memory requirements with the problem size, but, in this article, we focus on time complexity which investigates how the running time of the computation changes as the problem size increases.

Computational chemists often informally discuss commonplace concepts in computer science such as the complexity classes of polynomial-time problems (P) and non-deterministic polynomial-time problems (NP). For instance, it is sometimes stated that Hartree–Fock has a runtime which scales as the third power of the number of basis functions. This scaling disregards difficult instances of the calculation where Hartree–Fock does not converge. Such instances require manual intervention to tweak the algorithm used or adjust the convergence thresholds in a case-by-case fashion.

In computer science, often the focus is on worst-case complexity where the most difficult instances of a problem are used to classify the problem’s complexity. This is one of the major areas of theoretical computer science, although work on average case complexity does exist. In this article, the complexity of the problems discussed are characterized by the worst-case scaling.

2.1 Time complexity in equivalent computer models

A proper measure of the time complexity of an algorithm is how many basic operations (or how much time) it takes to solve problems of increasing size. Conventionally, a computational problem is described as easy or tractable if there exists an efficient algorithm for solving it, i.e. one that scales polynomially, $O(n^k)$ with fixed $k$ and input size $n$. Otherwise, the problem is considered intractable. This is an asymptotic definition that may not capture the full utility of the algorithm. For example, an asymptotically efficient algorithm may run slower than an asymptotically inefficient algorithm for small or fixed size problem instances. Nevertheless, this asymptotic classification of algorithms has proved useful. From a theoretical computer science perspective, the division allows for considerable progress to be made without considering the minutiae of the specific system, implementation, or domain of application.

From a practical standpoint, Moore’s law (or more aptly, Moore’s conjecture) states the density of transistors in classical computers doubles every two years. Thus, for a fixed computational time, if the computer cannot solve an instance when using a polynomial-time algorithm, one need not wait long as the exponential growth of computing power will reasonably quickly overtake the polynomially large cost. However, if the algorithm runs in exponential-time, one may be forced to wait several lifetimes in order for an instance to become solvable even if Moore’s law continues indefinitely. Complicating matters, the exponential growth according to Moore’s law is expected to cease sometime this century; hence the recent emphasis on quantum computation.

The time complexity can be characterized using any computationally equivalent model. In the context of computer science, equivalent means that one model can simulate the other with an overhead that scales polynomially as a function of system size.

† The notation $f(x) = O(g(x))$, implies that function $f(x)$ is bounded above by $g(x)$ times a constant for asymptotically large values of $x$ e.g. there exists a positive constant $C$ such that $Cg(x) \geq f(x)$ as $x \rightarrow \infty$. To indicated that $f(x)$ is asymptotically bounded below by $g(x)$, we write $f(x) = \Omega(g(x))$. If there are constants $C$ and $K$ such that $f(x)$ is bounded above by $Cg(x)$ and bounded below by $Kg(x)$ as $x$ goes to infinity, then we write $f(x) = \Theta(g(x))$. 

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These mappings respect the boundary between efficient and inefficient algorithms described earlier. Turing machines and circuits are two typical models discussed in the context of computational complexity.

The 'Turing computer' or 'Turing machine' was introduced by Alan Turing \(^{32}\) in 1936 before transistors and electronic circuits and formalizes the idea of a computer as a person to whom computational instruction could be given. Turing \(^{32}\) introduced the concept in order to answer David Hilbert's challenge to decide if a polynomial has roots which are integers using a "finite number of operations." Turing proved that this was not possible using his Turing machine. An illustration depicting the salient features of the Turing machine is given in Fig. 1.

The circuit model of computation has been more widely used when generalizing to the quantum setting although the quantum Turing machine formulation does exist. \(^{33,34}\) The time complexity in the circuit model is characterized by the number of circuit elements (or gates) used to compute the accept/reject boolean value corresponding to an input. So long as a universal set of gates, e.g. NAND, FAN-OUT, and FAN-IN for classical circuits, are used, the time complexity will only differ by a polynomial factor. The key caveat to consider when using this model is the notion of uniformity. To properly define an algorithm in the circuit model, a way of generating circuits for all input sizes must be given concisely. A circuit family for an algorithm is uniform if there is a polynomial-time algorithm that specifies the circuit given the input size.

2.2 Computational problems and problem instances

In computer science, computational problem often refers to decision problems which are collections of yes/no questions that an algorithm can decide. Each question is called a problem instance and the answer is called the solution to that instance.

Although we focus on decision problem, in many situations the decision problem can be used to extract numerical values by asking sufficiently many yes/no questions.

A collection of decision questions with affirmative answers such as, ‘‘Is 149 prime?’’ or ‘‘Is 79 prime?,’’ is called a language. Languages define decision problems where only the accept instances are included in the language. If an algorithm accepts (returns ‘‘True’’ or outputs 1) on all strings contained in a language, then the device is said to recognize the language. If the computer is also required to halt on all inputs, then it is said to decide the language. The difference being, a Turing machine may recognize a language but fail to halt on some instances, i.e. run forever. The problem of deciding if a Turing machine algorithm halts is called the HALTING problem. This problem has a rich and storied history beginning with the first paper of Turing. \(^{32}\) In Fig. 1, the algorithm decides the language of strings beginning with ‘‘01’’ or ‘‘10.’’

As an example from chemistry, the computational problem of deciding if a molecule has a dipole moment, \(D_{\text{POLE}}\), is a collection of questions: ‘‘Does \(x\) have a dipole moment?’’ Here, \(x\) is a string representing the molecule in each instance. The questions ‘‘Does \(\text{BCl}_3\) have a dipole moment?’’ and ‘‘Does \(\text{NH}_3\) have a dipole moment?’’ are instances of this computational problem. The string \(x = \"\text{BCl}_3\)’’ is not in the language \(D_{\text{POLE}}\) since ‘‘\(\text{BCl}_3\)’’ does not have dipole moment while ‘‘\(\text{NH}_3\)’’ is in the language since it has a dipole moment. A reasonable modification of the problem would be ‘‘Does \(x\) have a dipole moment greater than 0.1 Debyes?’’ so that small dipole moments may be ignored.

It may also be necessary to promise that \(x\) represents a molecule in a specific way such that ill formatted inputs can be ignored. This is accomplished using promise problems where the promise would be that \(x\) is indeed a string that properly encodes a molecule. Promise problems also arise in order to account for issues of numerical or experimental precision as illustrated in the next paragraph.

Promise problems also play a key role throughout the remainder of this text since we are discussing physical properties where infinite precision is neither required nor expected. As an illustration, we use an example from thermochemistry, where language \(A\) is the set of strings corresponding to (ideal) gases with heat capacity at constant pressure, \(C_p\), less than some critical value. Imagine you have an unreasonable lab instructor, who gives you a substance that has heat capacity extremely close to the critical value that decides membership in \(A\). It may take a large number of repetitions of the experimental protocol to be confident that the substance belongs or does not belong to language \(A\). A reasonable lab instructor would announce at the beginning of the lab that all the substances handed out for the experiment are promised to be at least one Joule per Kelvin away from the critical value. Given this promise, the student will be able to complete their lab in a single lab period. Without such a promise, it may take the student several lab periods to repeat the experiment in order to establish a sufficiently precise value of the heat capacity to decide the instance. Instead when using a promise, if the
student has a compound that violates the promise then the instructor would give full credit for either answer.

More formally, in the usual decision problems, the problem is defined using one language, \( L \), specifying the accept instances. The language of reject instances is all strings not in \( L \). Promise problems differ in that they are specified using two languages: one for the accept instances, \( L_{\text{accept}} \), and a separate language for the reject instances, \( L_{\text{reject}} \). If an instance is not in either of these languages, then the promise is violated and the computation can terminate with any result.

Reusing the \textsc{Dipole} example, if we can reformulate the problem to find out about non-zero dipole moments as a promise problem to account for experimental error. Now the computational task of \textsc{Dipole} is: ‘Given a molecule, \( x \), a trial value for the dipole \( d_T \), and an error tolerance \( \delta \), decide if the dipole moment of \( x \) is greater than \( d_T + \delta \) or less than \( d_T - \delta \), promised that the dipole moment is not between \( d_T \pm \delta \).’ In this case, there are two languages defining the problem \( L_{d_T=\delta} \) and \( L_{d<d_T-\delta} \). If a molecule has a dipole moment of exactly \( d_T \), it would violate the promise and the experimenter or computer does not need to answer or can respond with any answer. The value \( \delta \) allows us to meaningfully define problems in the presence of errors resulting from imperfect experimental measurements or from the finite precision of a computing device.

2.3 Computational reductions

The idea of computational reduction is at the heart of classifying computational problems. Reducibility is a way of formalizing the relationship between problems. Essentially, it asks, “If I can solve one problem, what other problems can I solve using the same resources?”

There are two main types of reductions: Turing reductions and Karp reductions. For the reductions to be useful, they must be limited to polynomial time. The polynomial-time Turing reduction, also called the Cook reduction, of problem \( A \) to problem \( B \) uses solutions to multiple instances of \( B \) to decide the solutions for instances of \( A \). The solutions to instances of \( B \) are provided by an oracle and each questions is called a query. Algorithms deciding \( A \) which require only polynomial queries of the oracle for \( B \) are efficient whenever the oracle for \( B \) answers in polynomial time.

The other type of reduction is called a Karp reduction or polynomial transformation. If, instead of an oracle, there is an embedding of instances of problem \( A \) into instances of problem \( B \), then an efficient solution for all instances of problem \( B \) imply efficient solutions for instances of problem \( A \). This transformation is a Karp reduction. When \( A \) can be reduced to \( B \), under either Karp or Turing reductions, it is denoted \( A \leq B \).

To illustrate the difference, we use examples from thermodynamics. Consider trying to determine if the heat capacity at constant pressure, \( C_p \), of an ideal substance is less than some critical value (language \( A \)). Now, suppose an oracle, probably in the form of a bomb calorimeter, has been provided for deciding if the heat capacity at constant volume, \( C_V \), of a given substance is above or below a critical value. Call language \( B \) the set of strings labeling substances with \( C_V \) below this value. By adding the number of moles times the ideal gas constant to the critical value, one can determine membership in language \( A \) via the formula \( C_p = C_V + nR \).

Since each instance of \( A \) can be embedded into an instance of \( B \), language \( A \) is Karp reducible to \( B \) and we can write \( A \leq B \).

Suppose instead, an oracle for evaluating if, at fixed pressure, the internal energy, \( U \), of the given substance at a given temperature is less than some critical value (language \( C \)) is given. Then by evaluating the internal energy at two different temperatures, the heat capacity at constant pressure can be bounded by numerically estimating the derivative. Because the reduction has to use the oracle for \( C \) more than once to decide instances of \( A \), language \( A \) is Turing reducible to \( C \) and \( A \leq C \).

2.4 Basic complexity classes

Equipped with the basic definitions from computer science, we now introduce the concept of computational complexity classes. This will give some insights into why quantum chemistry is difficult. We will introduce six basic complexity classes to classify the time complexity of decision problems. The first three complexity classes are characterized by algorithms that can decide instances in time proportional to a polynomial of the input size; the other classes are characterized by polynomial time verification of problem instances.

In Table 1, the three polynomial time complexity classes are listed. First, \( P \), is the class of all decision problems with instances that can be accepted or rejected by a Turing machine in polynomial time. If the Turing machine has access to an unbiased random number generator, then the complexity class of problems that can be decided in polynomial time is called BPP. The term “bounded error” refers to the requirement that the probability of acceptance and of rejection must be bounded away from half so that repetition

<table>
<thead>
<tr>
<th>Class</th>
<th>Name</th>
<th>Computer type taking only poly. time</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>Polynomial time</td>
<td>Turing machine</td>
</tr>
<tr>
<td>BPP</td>
<td>Bounded error probabilistic polynomial time</td>
<td>Turing machine with access to random number generator</td>
</tr>
<tr>
<td>BQP</td>
<td>Bounded error quantum polynomial time</td>
<td>Turing machine with access to quantum resources</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Class</th>
<th>Name</th>
<th>Verifier’s computer type taking only poly. time</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP</td>
<td>Non-deterministic polynomial time</td>
<td>Turing machine</td>
</tr>
<tr>
<td>MA</td>
<td>Merlin–Arthur</td>
<td>Turing machine with access to random number generator</td>
</tr>
<tr>
<td>QMA</td>
<td>Quantum Merlin–Arthur</td>
<td>Turing machine with access to quantum resources</td>
</tr>
</tbody>
</table>

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can be employed to boost the confidence in the answer. An
important class of problems falling into this complexity class are
efficient Monte Carlo simulations often grouped under the umbrella
term quantum Monte Carlo \cite{55-37} used for computing electronic
structure in chemistry. Lastly, the complexity class BQP is character-
ized by problems solvable in polynomial time with quantum
resources. In the quantum computational model, \cite{38} the quantum
algorithm is conceptually simpler to think of as a unitary circuit, \( U \),
composed of unitary circuit element that affect, at most, two
quantum bits. As mentioned earlier, the number of gates used
determines the time complexity. The outcome of the algorithm with
input \([\text{Input}]\) is “accept” with probability \([\langle \text{Accept} \rangle(U)\text{Input}]\)^2. Similarly,
for the “reject” cases.

So far, the discussion has centered on complexity classes con-
taining problems that are solvable in polynomial time given access
to various resources: a computer (P), a random number generator
(BPP), and a controllable quantum system (BQP). Now, our attention
turns to the class of problems that can be computed non-deter-
ministically. The original notion of non-determinism is a Turing
machine whose transition rule maps the computer state and the
tape symbol to any number of possible outputs. In essence, this is a
computer that can clone itself at will to pursue all options at once.
NP is the class of problems that could be solved in polynomial time
by such a computer. Whether a deterministic polynomial time
computer can be used to simulate such a computer is a restatement
of the famous open question in computer science: “Does P = NP?”
This question was selected as a millennium problem by the Clay
Mathematics Institute which has offered a one million dollar prize
for a correct proof of the answer.

Rather than resorting to imaginary computers, the non-
deterministic classes can be defined by considering verification
of solutions which have been obtained in some non-determin-
istic way, see Fig. 2. For every “accept” instance, \( x \), there exists
at least one proof state \( y \) such that the verifier returns “accept”
in polynomial time. If \( x \) is should be rejected, then for every
proof state \( y \), the verifier should output “reject”, again, in a
polynomial amount of time. Each of the non-deterministic classes
listed in Table 1, are characterized by the computational
dpower of the verifier. Note that P is a subset of NP because any
problem that can be easily solved can be easily verified.

2.5 Completeness and hardness
The goal of classifying computational problems into complexity
classes motivates introduction of the terms hard and complete.
The classification of a computational problems as hard for a
complexity class means if an algorithm can solve this problem,
then all problems in the class can be solved efficiently. In other
words, this problem contains all problems in the class either
via Karp or Turing reductions. More precisely, a language \( B \) is
hard for a complexity class \( CC \) if every language, \( A \), in \( CC \) is
polynomial-time reducible to \( B \), i.e. \( A \leq B \).

Complete problems for a class are, in some sense, the hardest
problems in that class. These problems are both a member of
the class and are also hard for the class. That is, a language \( B \) is
complete for a complexity class \( CC \) if \( B \) is in the complexity class
\( CC \) and for every other language \( A \) in \( CC \), \( A \leq B \).

The simplest illustration of the difference between hard and
complete computational problems is the distinction between opti-
mization problems and decision problems. Optimization problems
such as finding saddle points or minima in energy landscapes are
frequently encountered in computational chemistry, but these
problems are not in the complexity class NP. However, if one can
perform optimization quickly, then responding with the answer to
yes/no questions about the solution would be simple. Thus,
optimization problems can be classified as NP-hard but not
NP-complete since the computational task of optimization is, in a
sense, harder than just answering yes or no.

3 Electronic structure and other Hamiltonian
problems
Now, armed with the key ideas from computer science, we return
to our original inquiry into why quantum chemistry is hard. We
answer this questions by exploring the computational complexity
of three different widely used methods for computing electronic
energy in computational chemistry: Hartree–Fock, two-electron
reduced density matrix methods, and density functional theory.

Before delving into the specific complexities of these
problems, we first establish notation by reviewing necessary
concepts from quantum chemistry, then discussing computa-
tional problems concerning classical and quantum spin Hamil-
tonians. We end the section with mappings between systems
of electrons and spin systems.

3.1 Quantum chemistry and second quantization
In quantum chemistry, the annihilation, \( \{a_i\} \), and creation
operators, \( \{d_i\} \), correspond respectively to removing and adding
an electron into one of \( M \) single particle wave functions, \( \{\phi_i(x)\}_{x=1}^M \).
The single-particle wave functions are called orbitals and the set
of orbitals is typically called the basis set. To include the electron spin,
the single particle orbitals \( \phi_i \) are functions of spatial coordinates
and a spin variable which are collectively denoted \( x \).

The electronic spin will play an important role when dis-
cussing the connections between systems of electrons and
systems of quantum spins. Since electrons are spin-\( 1/2 \) particles,
the electron spin is either up or down. Accordingly, there are
\( M/2 \) orbitals with spin up and \( M/2 \) with spin down. Unless
explicitly noted, the summation over orbitals includes a sum-
mation over the spatial and spin indices of the orbitals.

Anti-symmetry of the \( N \)-electron is enforced by the canonical
anti-commutation relations for fermions, i.e. electrons,
\[ [a_p, a_q]^\dagger = a_p^\dagger a_q + a_q^\dagger a_p = 0, \quad [a_p, a_q^\dagger] = \delta_{pq} 1. \] (1)

In chemistry, typically the electronic structure is often the
primary concern, but if interested in vibrational structure, we
would have to consider bosonic canonical commutation rela-
tions: \[ [b_p, b_q] = b_p^\dagger b_q - b_q b_p^\dagger = 0 \text{ and } [b_p, b_q^\dagger] = \delta_{pq} 1. \] The vacuum
state, \( |\text{vac} \rangle \), is the normalized state that the creation/annihil-
ation operators act on and it represents the system with no
particles.
Acting on the vacuum state with a strings of $N$ distinct creation operators yields $N$-electron wave functions. The most general $N$-electron state within a basis set is,

$$|\Psi\rangle = \sum_k C_k a_{k1}^\dagger a_{k2}^\dagger \cdots a_{kn}^\dagger |\text{vac}\rangle,$$

(2)

with $K = (K_1, \ldots, K_N)$ and the complex valued $C_k$ are constrained such that $\sum_k |C_k|^2 = 1$. To convert between creation and annihilation operators and the coordinate representation, one uses field operators: $\hat{\phi}(x) = \sum_k \phi_k(x)a_k$ as in $\Psi(x_1, \ldots, x_N) = \langle \text{vac}|\hat{\phi}(x_1)\cdots\hat{\phi}(x_N)|\Psi\rangle/\sqrt{N!}$. The factor of $N!$ accounts for permutation symmetries in the summation. Each state $\Psi$ or $|\Psi\rangle$ is called an $N$-electron pure state. A valid $N$-electron mixed state,

$$\rho^{(N)} = \sum_i p_i |\Psi_i\rangle\langle\Psi_i|$$

(3)

has $\sum p_i = 1$ where each $|\Psi_i\rangle$ is an $N$-electron wave function. The $k$-th order reduced density matrix, abbreviated $k$-RDM, is defined using the field operators:

$$\rho^{(k)}(x_1, x_1', \ldots, x_k, x_k') = \frac{1}{(N)_k} \langle \Psi|\hat{\phi}^\dagger(x_1)\cdots\hat{\phi}^\dagger(x_k)\hat{\phi}(x_1')\cdots\hat{\phi}(x_k')|\Psi\rangle$$

(4)

$$= \frac{1}{(N)_k} \text{Tr} \left[ \hat{\phi}(x_1)\cdots\hat{\phi}(x_k) \rho^{(N)} \hat{\phi}^\dagger(x_1)\cdots\hat{\phi}^\dagger(x_k) \right]$$

(5)

With $(N)_k = N!/(N-k)!$, the reduced density matrices are normalized to unity and, in eqn (5), the trace sums over the expectation values of states from a complete set of $(N-k)$-electron wave functions.

Lastly, we define the primary computational chemistry problem encountered in computation of electronic structure:

**Electronic Structure.** The inputs are the number of electrons, $N$, a set of $M$ orbitals, a static configuration of nuclei, a trial energy, $E_T$, an error tolerance $\delta$ and Hamiltonian,

$$H_{\text{elec}} = T_e + V_{ee} + V_{en} = \sum_{ij} (T_{ij}^e + V_{ij}^{(N)}) a_i^\dagger a_j + V_{ij}^{ee} a_i^\dagger a_i a_j a_j$$

(6)

with $T_e$ the electronic kinetic energy operator, $V_{ee}$ the electron-electron interaction operator and $V_{en}$ the electron-nuclear interaction. The task is to decide if the ground state energy is less than $E_T - \delta$ or greater than $E_T + \delta$ promised that the energy is not between $E_T \pm \delta$.

Methods related to getting approximate solutions to **Electronic Structure** have complexities ranging from NP-complete to QMA-complete as shown in Sections 4 and 5. The complexity of **Electronic Structure** itself is not yet clear. In the literature, only Hamiltonians with more flexibility, namely a local magnetic field, have been demonstrated to be QMA-complete as shown see Section 6.1.

### 3.2 Classical and quantum spin Hamiltonians

The problem of estimating the ground-state energy of Hamiltonians of different forms lies at the intersection of computer science, physics and chemistry. In this section, we define computational problems related to both classical and quantum spin Hamiltonians. Spin systems have long been known to provide fertile ground for complexity theory.\textsuperscript{19,39–45} Given a Hamiltonian, deciding if there exists a spin configuration, e.g. $\{1 \uparrow \downarrow \cdots \}$, that satisfies a certain property, e.g. has energy below a certain threshold, can be difficult even when checking the property for each configuration is easy. Computations designed to decide properties of this sort are often in the complexity class NP. If, additionally, that particular property can embed instances of any other NP properties, then that property corresponds to an NP-complete computational problem. A canonical example is the following spin problem:

**Ising.** The inputs are $N$ classical spins with possible values $\pm 1$, trial energy, $E_T$, and Ising Hamiltonian

$$H_{\text{ising}} = - \sum_{\langle ij \rangle} J_{ij} S_i S_j$$

(7)

having $J_{ij}$ either 0 or $\pm 1$. The task is to decide if there is a configuration of spins $c = (s_1, s_2, \ldots, s_N)$ such that the energy of the Ising Hamiltonian is less than $E_T$ or if all configurations have energy above $E_T$.

Unlike the other Hamiltonian problems discussed in this review, Ising does not require a promise on the precision.
because the energy values of the Hamiltonian are integer-valued instead of continuous.

Although some instances of the Ising lattice can be solved analytically, e.g. one-dimensional Ising lattices or the solution by Onsager\textsuperscript{42} for two-dimensional Ising lattices with uniform couplings, Barahona\textsuperscript{39} showed that problem Ising is NP-hard when interactions are restricted to nearest neighbors on a $2 \times L \times L$ lattice. Since this model is only of tangential interest to electronic structure, details of this proof are omitted. Since the “accept” instances can be verified in polynomial time given a configuration $c$ with energy less than $E_D$, this problem is in the complexity class NP and hence, is NP-complete. Further results show that all non-planar lattices, even in two-dimensions, are NP-complete.\textsuperscript{40}

To provide examples of Hamiltonian problems that are complete for the quantum analogue of NP, we turn to the quantum analogues of the Ising Hamiltonian: quantum spin Hamiltonians. To define similar quantum Hamiltonians with QMA-complete properties, it turns out that one only needs spins with angular momentum of $\frac{1}{2}$ which are often called 	extit{qubits}. The interactions are now expressed using tensor products of the Pauli sigma matrices with matrix representations:

\[
\sigma^x = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}, \quad \sigma^y = \begin{bmatrix} 0 & -i \\ i & 0 \end{bmatrix}, \quad \sigma^z = \begin{bmatrix} 1 & 0 \\ 0 & -1 \end{bmatrix}.
\]

Additionally, $\sigma^0$ is the identity matrix.

A widely studied class of qubit Hamiltonian problems whose computational complexity was first investigated by Kitaev et al.\textsuperscript{41} is the problem of computing, to polynomial accuracy, the ground state energy of $k$-spin local Hamiltonians. This set of problems is referred to as $k$-\textit{LOCAL HAMILTONIAN} in the computational complexity literature following ref. 41 but we use the name $k$-\textit{LOCAL SPIN HAMILTONIAN} to emphasize the nature of the Hamiltonian. That is, $k$-\textit{LOCAL SPIN HAMILTONIAN}. Given $k$-spin-local Hamiltonian acting on $N$ spins,

\[
H_{\text{QMA}} = \sum_{D = 0}^{m} \sum_{C = 0}^{\text{poly}(N)} J_{C,D} \sigma^d_{c_1} \otimes \sigma^d_{c_2} \otimes \cdots \otimes \sigma^d_{c_k}\tag{9}
\]

where $d_i \in \{x, y, z, 0\}$, $c_i$ labels a spin, there are $m = \text{poly}(N)$ terms, and $|J_{C,D}| \leq 1$, decide if the ground state energy is less that $E_0 - \delta$ or if the ground state energy is greater than $E_0 + \delta$ with $\delta < 1/\text{poly}(N)$ promised that is not between $E_0 \pm \delta$.

Note that the problem is defined so that the error scales relative to the number of non-zero terms $m$ in eqn (9). To understand this, consider the following set of instances with error tolerance $\delta = 0.5$. Consider Hamiltonian, $H$, with ground state energy 17.7 and Hamiltonian, $H'$, with energy 17.6. As it stands, the two Hamiltonians cannot be distinguished. However, by including each term in $H$ and $H'$ ten additional times, the ground state energies are now 177 and 176, which can be resolved at error tolerance $\delta$. By considering $m$ as a polynomial in $N$, one cannot indefinitely rescale the energy to effectively shrink the error tolerance.

The first demonstration of a QMA-complete problem\textsuperscript{41} required $k = 5$. Subsequently $k$ was reduced\textsuperscript{43} to 2. Finally, the problem was shown\textsuperscript{44} to remain QMA-complete even when $J_{C,D}$ is non-zero only if two spins are spatially adjacent on a 2D lattice. The problem can be further restricted by including only a limited set of two-spin interactions and still remain QMA-complete.\textsuperscript{45} Complexity results concerning qubit Hamiltonians as well as other variants with higher dimensional spins and more restrictive lattices were recently reviewed by Osborne.\textsuperscript{19}

### 3.3 Relationships between spin systems and electronic systems

Since chemists are not necessarily interested in qubit or quantum spin systems, we now discuss connections to systems of indistinguishable electrons. In this subsection, we will examine how to embed a system of spins into an electronic Hamiltonian and how to embed electronic systems into spin Hamiltonians. Lastly, we use these connections to give an example of a fermionic system (although not Electronic Structure) that is QMA-complete.

Given an electronic Hamiltonian, there is a \textit{orbital pair pseudo-spin} mapping\textsuperscript{46,47} that is used to embed qubit models to the ground state of the electronic systems. To embed $N$ spins, there must be $M = 2N$ orbitals and $N$ electrons, i.e. half-filling, and each quantum spin is identified with a pair of fermionic modes. The embedding requires translating each spin $|\sigma> = |\uparrow> + |\downarrow>$ to fermionic operators: $\alpha|\uparrow> + \beta|\downarrow>$. The Pauli operators appearing in the spin Hamiltonian, then become single fermion terms, e.g. $h_{ij}a_i^\dagger a_j$. As important examples consider,

\[
\sigma^x_i = |\downarrow> \langle \downarrow| + |\uparrow> \langle \uparrow| \\
\sigma^y_i = i |\downarrow> \langle \uparrow| - |\uparrow> \langle \downarrow| \\
\sigma^z_i = |\uparrow> \langle \uparrow| - |\downarrow> \langle \downarrow|
\]

Taking tensor products of the Pauli matrices leads to two-fermion terms, e.g. $h_{\pi}a_i^\dagger a_j^\dagger a_j a_i$. The final concern is preventing double occupancy within a pair of sites which would invalidate the pseudo-spin interpretation. This is handled\textsuperscript{47–49} by including an additional term which penalizes invalid electronic configurations. For each pair of electronic modes, the two-fermion penalty

\[
P_i = C|\alpha|^2|a_i^\dagger a_j|^2(11)
\]

is added to the electronic Hamiltonian. Because penalty $P = \sum P_i$ commutes with the fermionic Pauli matrices, the ground state still corresponds to the solution of the spin Hamiltonian. The constant $C$ can be selected as a low order polynomial in the system size to ensure that the ground state remains in a valid pseudo-spin state.\textsuperscript{‡} Note that we have not explicitly relied on the anti-commutation properties and a nearly identical construction exists for bosonic systems.\textsuperscript{50}

A second connection is given using well established techniques developed to translate certain spin systems to simpler non-interacting fermionic systems that can be exactly solved.\textsuperscript{51–53} The Jordan–Wigner transform\textsuperscript{54} provides this connection by mapping fermions to spin operators such that the canonical anti-
commutation relations are preserved. The Jordan–Wigner transform between \( N \) fermionic creation and annihilation operators and the Pauli matrices acting on \( N \) spins is given by

\[
a_j \leftrightarrow 1^{l_{j-1}} \otimes \sigma^+ \otimes \sigma^z^{l_{N-j}} \quad (12a)
\]

\[
a_j \dagger \leftrightarrow 1^{l_{j-1}} \otimes \sigma^- \otimes \sigma^z^{l_{N-j}} \quad (12b)
\]

where \( \sigma^+ = \frac{1}{\sqrt{2}} (| \uparrow \rangle \langle \downarrow | + | \downarrow \rangle \langle \uparrow |) \) and \( \sigma^- = \frac{1}{\sqrt{2}} (| \downarrow \rangle \langle \uparrow | - | \uparrow \rangle \langle \downarrow |) \). The qubit state \( | \downarrow \downarrow \rangle \) corresponds to the vacuum state and the string of \( \sigma^z \) operators, preserve the commutation relations in eqn (1) since \( \sigma^+ \) and \( \sigma^- \) anti-commute. More sophisticated generalizations of the Jordan–Wigner transform, reduce the number and the spatial extent of the spin–spin interactions.\(^{54-56}\)

The orbital pair pseudo-spin mapping and the Jordan–Wigner transformation complement each other and will be used repeatedly throughout the remainder of the article. The orbital pair pseudo-spin mapping requires carefully engineered penalties and fixes the total number of orbitals. Thus, it is primarily useful when translating spin systems to electronic systems. By contrast, the Jordan–Wigner transformation is primarily useful in the other direction, that is, when translating an arbitrary electronic system to a quantum spin system. The next subsection gives a concrete example of how these connections are employed when studying the computational complexity of electronic systems.

### 3.4 Generic local fermionic problems are QMA-complete

The orbital pair construction allows one to immediately ascertain that the ground state energy decision problem for Hamiltonians containing two-fermion interactions,

\[
H_{sf} = \sum_{j} h_{j} a_{j} a_{j}^\dagger + h_{jk} a_{j} a_{k} a_{j} a_{k}^\dagger
\]

is QMA-hard.\(^{48}\) This follows via a Karp reduction using the orbital pair pseudo-spins as in eqn (10). Since only two-fermion interactions in eqn (9) are required for QMA-completeness,\(^{13}\) each spin–spin interaction term translates to two-fermion terms under the pseudo-spin mapping. Since the pseudo-spin construction can be extended to bosonic systems, the two-boson ground state energy decision problem is also QMA-hard.\(^{50}\)

The ground state energy of the electronic state can be verified to be above or below \( E_T \pm \delta \) by a BQP quantum computer given the proof state. This implies that the problem is QMA-complete. We sketch the idea relying on well known results about BQP quantum simulation of chemical systems.\(^{5,6,7}\) The Jordan–Wigner transformation is used to translate the two-electron Hamiltonian into a qubit Hamiltonian. If Merlin provides the fermionic ground state,\(^8\) the energy of corresponding qubit state can be determined through simulation\(^{8,57,58}\) of the evolution under the qubit version of the two-electron Hamiltonian. This quantum evolution can be simulated efficiently on a quantum computer.\(^{59,60}\) The resulting evolution in the time-domain is Fourier transformed to extract the energy of the ground state\(^{5,6,11,61}\) allowing the verifier to determine whether to accept or reject the instance. This argument carries through for the bosonic case as well using the bosonic equivalent of the Jordan–Wigner transform.\(^{50}\)

### 4 Hartree–Fock

Hartree–Fock is one of the most important quantum chemistry techniques as it typically recovers about 99% of the total electronic energy. Hartree–Fock is known to be a weak approximation in many instances, but it is the basis for more sophisticated (post-Hartree–Fock) methods which improve upon the Hartree–Fock wave function. Furthermore, Hartree–Fock provides the mathematical framework for the widely adopted notion of molecular orbitals used throughout chemistry.

The implementation of the Hartree–Fock algorithm requires diagonalizing the \( M \) by \( M \) Fock matrix at \( O(M^3) \) cost. When this dominates the runtime, the algorithm then scales as a third-order polynomial in the size of the basis. However, since Hartree–Fock solves a nonlinear eigenvalue equation through an iterative method,\(^1\) the convergence of self consistent implementations is the key obstacle that prevents the worst-case scaling from being polynomial. While many rank and file computational packages are routinely employed for self consistent field calculations, there are well documented molecular instances involving actinides or transition metals where convergence is known to be a problem. The following complexity result highlights the fact that convergence problems are intrinsic to the difficulty of the problem being solved. The computational complexity result proving that the worst-case scaling cannot be polynomial unless \( \text{P=NP} \) was provided in an unpublished appendix of Schuch and Verstraete\(^{49}\) available on the arXiv preprint server.

For the purposes of this article, the Hartree–Fock procedure can be succinctly explained as the minimization of the energy of an \( N \)-electron system given \( M \) basis functions with the restriction that in the expansion found in eqn (2), all \( C_k \) are zero except one. Explicitly,

\[
E_{HF} = \min_{\Psi} \sum_{k=1}^{N} \langle \Psi | H_{dec} | \Psi \rangle
\]

Here, the single Fock state corresponding to the minimal value of \( E_{HF} \) is called the Hartree–Fock state: \( \Psi_{HF} \). The optimized set of creation operators,

\[
h_j^{\dagger} = \sum_{j} C_{j} a_j^{\dagger}
\]

place electrons into the set of molecular orbitals, \( \psi_{j}(x) = \sum_{i} C_{ij} \phi_{j}(x) \). The formal computational problem \( \text{HARTREE–FOCK} \) will be defined analogous to the other Hamiltonian problems with a promise given to account for precision. Specifically,

\( \text{HARTREE–FOCK} \). The inputs are the number of electrons, \( N \), a set of \( M \) orbitals, a trial energy, \( E_T \), an error tolerance, \( \delta < 1/\text{poly}(N) \), and a two-electron Hamiltonian, \( \text{cf. eqn (13)} \). The sum of the absolute values of the coefficients is required to

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\( \text{cf. eqn (13)} \). The sum of the absolute values of the coefficients is required to
scale less than a polynomial in $N$. The task is to decide if the Hartree–Fock energy $E_{HF} = \langle \Psi_{HF} | H | \Psi_{HF} \rangle$ is less than $E_T - \delta$ or greater than $E_T + \delta$ promised that $E_{HF}$ is not between $E_T \pm \delta$.

### 4.1 Hartree–Fock is NP-complete

To prove that any other NP problem can be mapped to the HF problem (NP-hardness), we can use a Karp reduction to embed instances of Ising into instances of HF.\(^{49}\) Using the fermionic version of $\sigma^2$ given in eqn (10), $H_{\text{Ising}}$ only requires two-electron interactions,

$$S_i S_j = \sigma_i^x \sigma_j^x = (a_i \dagger a_i - a_i^a_i)(a_j \dagger a_j - a_j^a_j)$$

The satisfying assignment of the Ising instance is some spin configuration $|s_1, s_2, \ldots, s_N\rangle$ for $N = 2L^2$ with $s_i$ as either $\uparrow$ or $\downarrow$. The correct and exact Hartree–Fock solution for the $N$-electron wave function should assign $b_i = a_i^\dagger$ when $s_i = \uparrow$ and $b_i = a_i$ when $s_i = \downarrow$. Thus, $\text{Ising} \leq \text{Hartree–Fock}$ under a Karp reduction.

The promise on the error tolerance, $\delta$, given in the specification of the problem is necessary because of the pseudo-spin representation used in eqn (16). Corrections to the energy due to the pseudo-spin representation can be computed using perturbation theory with the unperturbed Hamiltonian given in eqn (11) and the converted Ising Hamiltonian, cf. eqn (16), as, the perturbation. The first order corrections in small parameter $C^{-1}$ are the Ising energies and the errors due to the pseudo-spin mapping arise at second order in $C^{-1}$. A coarse estimate for $C$ is obtained by multiplying the number of non-zero terms by the maximum absolute value of a coefficient occurring in eqn (13). Since we map the problem to Ising, there are $O(N^2)$ terms in the summation and the maximum absolute value of each term is unity. Hence, $C$ is estimated as $O(N^2)$. So long as $\delta < O(N^{-2})$, the first order corrections occurring at order $C^{-1}$ can be distinguished and the ground state Ising energies can be recovered.

To prove inclusion of HF in the complexity class NP, an algorithm for verifying the energy in polynomial time must be given. If the coefficient matrix $C$ describing the correct orbital rotation is given, then the energy is calculated easily using Slater–Condon rules.\(^{62}\) Thus, given the answer, the accept/reject conditions are quickly verified. Since the problem can also be quickly verified, Hartree–Fock is NP-complete.

### 5 2-RDM methods

Many computational chemistry algorithms seek to minimize the energy or other properties by manipulating the wave function but to quote Coulson,\(^{63}\) ‘wave functions tell us more than we need to know . . ..’ All the necessary information required for energy and calculating properties of molecules is embodied in the first and second order density matrices’. Extensive work has been done to transform this remark into a host of computational methods in quantum chemistry.\(^{64,65}\) In this section and the following, we review the prominent computational complexity results related to this body of work.

With respect to the 2-RDMs, it is easy to evaluate properties of the electronic system since the Hamiltonian only contains two-electron terms. However, difficulties arise when determining if the 2-RDM is valid or invalid. Unconstrained optimization of the energy using 2-RDM can lead to ground state energies of negative infinity if the validity of each 2-RDM cannot be determined. While the criteria for validity have recently been developed\(^{66,67}\) deciding the validity of each 2-RDM using such criteria is known as the $N$-representability problem. Mazziotti\(^{66}\) also provides a proof that the following problem, $N$-Rep, is at least NP-hard. In the next subsection, we follow Liu et al.\(^{48}\) to demonstrate the stronger conclusion that $N$-Rep is QMA-complete.

#### 5.1 $N$-Rep is QMA-complete

To show QMA-hardness, a Turing reduction is used to show $\text{2-LOCAL-SPIN HAMILTONIAN} \leq \text{N-Rep}$. Before proceeding to the Turing reduction, note, since $N$-Rep only considers a fixed number of electrons, the one-electron operators are not needed as $\sum_{i \neq j} a_i^\dagger a_j / (N - 1)$. Each valid 2-RDM can be represented by an $M \times M$ dimensional matrix with $\binom{M}{2}$ independent parameters.\(^{\dag}\) A complete set of observables, such as the projection onto each matrix element,\(^{**}\) is then used to characterize the space of 2-RDMs.

The space of valid 2-RDMs is convex; that is if $\rho^{(2)}_1, \rho^{(2)}_2, \ldots, \rho^{(2)}_t$ are valid 2-RDMs, then the convex sum $\sum_{j} \rho^{(2)}_j$ with $\sum \rho^{(2)}_j = 1$ is also a valid 2-RDM. This follows directly from the convexity properties of sets of density matrices and probability distributions.

With access to an oracle for $N$-Rep, the boundaries of the convex set of valid 2-RDMs can be characterized. Because the oracle is used multiple times throughout the verification procedure, this is a Turing reduction. Since convex minimization problems can be solved efficiently and reliably, see e.g.,\(^{68}\) the QMA-hardness is nearly demonstrated. The last point of concern addressed by Liu et al.,\(^{48}\) are the errors introduced by the promise given on the oracle. To demonstrate that the algorithm remains robust in spite of such errors, the authors use a tailored version of the shallow-cut ellipsoid convex optimization technique.\(^{69}\)

\(\dag\) The appropriate metric is the trace distance, $d_{tr}(A, B) = \| A - B \|_1 = \frac{1}{2} \text{Tr} \left( \sqrt{(A - B)^\dagger (A - B)} \right)$, which generalizes the distance metric from standard probability theory.

\(\ddagger\) Since each 2-RDM is hermitian, the complex off-diagonal elements occur in pairs and the diagonal elements must be real. Since the trace is normalized to unity, there a reduction of one degree of freedom, leaving $\binom{M}{2} - 1$ independent parameters.

\(\ddagger\) Liu et al.\(^{48}\) used a different set observables inspired by the Pauli matrices with more convenient mathematical properties.
Let us remark that the required error tolerance follows since the reduction relies on the QMA-completeness of 2-\textsc{local-spin Hamiltonian} which includes a promise on allowed the error tolerance.

To demonstrate that \( N\text{-Rep} \) is QMA-complete, what remains is demonstrating that \( N\text{-Rep} \) is in the complexity class QMA. If \( \mu^{(2)} \) is a valid 2-RDM, then, relying on the Jordan–Wigner transform introduced earlier, Merlin can send polynomial copies of the correct \( N \)-electron state, \( \mu^{(N)} \). The verifier, Arthur, first checks that the number of electrons in the given state is \( N \) by evaluating \( \text{Tr}[\mu^{(N)} \sum a_i^\dagger a_k] \). Then Arthur randomly picks observables from the complete set described earlier and tests that the expectation value of the state \( \mu^{(N)} \) and \( \mu^{(2)} \) match until convinced. Merlin might try to cheat by sending entangled copies of \( \mu^{(N)} \) but it can be proven that Arthur cannot be fooled based on the Markov inequality.†† See Aharonov and Regev70 for a proof of this fact.

5.2 Restriction to pure states

The pure state restriction of \( N\text{-Rep} \) has interesting consequences. Consider, pure \( N\text{-Rep} \), where the question is now: “Did the 2-RDM come from a \( N \)-electron pure state (up to error \( \beta \))?”. This contrasts with the original problem where the consistency questions refers to an \( N \)-electron mixed state.

This problem is no longer in QMA since the verifier, Arthur, cannot easily check that the state is pure if the prover, Merlin, cheats by sending entangled copies of \( |\psi^{(N)}\rangle \). If instead, two independent unentangled provers send the proof state, the state from the first prover can be used to verify the purity. The purity is checked using the pairwise swap test,71 a quantum algorithmic version of the Hong–Ou–Mandel effect in quantum optics,72 on each of the supposedly unentangled states. When the given state is not a product state (or nearly so), Arthur will detect it. If the set of states from the first prover pass the test, the second set can be used to verify that randomly selected expectation values of \( |\psi^{(N)}\rangle \) and \( \mu^{(2)} \) match. The complexity class where two provers are used instead of one is QMA(2).

6 Density functional theory

In this section, a further reduced description is examined: density functional theory. Density functional theory (DFT) methods are of profound importance in computational chemistry due to their speed and reasonable accuracy.6,5,6,5,73 In this section, the complexity of the difficult aspects of DFT is shown to be QMA-complete. This was first conjectured by Rassolov and Garashchuk20 and rigorously proven by Schuch and Verstraete.49

In DFT, the wave function is replaced by the one-electron probability density, \( n(x) = \mu^{(1)}(x,x) \). The use of an object in three spatial dimensions to replace an object in \( 3N \) spatial dimensions without losing any information seems almost absurd, but the theoretical foundations of DFT are well established.65,74 The Hohenberg–Kohn theorem74 proves that the probability density obtained from the ground state wave function of electrons in a molecular system is in one-to-one correspondence with the external potential usually arising from the static nuclear charges. Therefore, all properties of the system are determined by the one-electron probability density.

In the proof of the Hohenberg–Kohn theorem, one of the most important, and, elusive, functionals of the one-electron density is encountered: the universal functional. In all electronic Hamiltonians, the electrons possess kinetic energy and the electrons interact via the Coulomb interaction; only \( V_{\text{ex}} \) due to the nuclear configuration changes from system to system. Separating these parts allows one to define the universal functional of DFT as

\[
F[n(x)] = \min_{\mu^{(1)}(x,x)} \text{Tr}[(T_e + V_{\text{ee}})\mu^{(N)}],
\]

which takes as input the probability density and returns the lowest possible energy of \( T_e + V_{\text{ee}} \) consistent with the probability density. The nuclear potential energy can be directly determined efficiently using \( \int dx \, n(x) V_{\text{ex}}(x) \) using Gaussian quadratures or Monte Carlo sampling. Regardless of the method used to evaluate the integral, the domain does not (explicitly) increase with the number of electrons. As a decision problem, we have the following computational problem:

\textbf{Universal Functional.} The inputs are the number of electrons, \( N \), a probability density \( n(x) \), a trial energy \( E_T \) and an error tolerance \( \delta < 1/\text{poly}(N) \). The task is to decide if \( F[n(x)] \) is greater than \( E_T + \delta \) or less than \( E_T - \delta \) promised that \( F[n(x)] \) is not between \( E_T - \delta \). It is required that the summation over the Hamiltonian coefficients \( \sum |V_{\text{ee}}^{\text{ex}}| + |T_e^2| + |V_{\text{ee}}^{\text{mag}}| + |V_{\text{mag}}^{\text{mag}}| \) scale less than \( \text{poly}(N) \).

6.1 Universal Functional is QMA-complete

The demonstration that \textbf{Universal Functional} is QMA-hard proceeds \textit{via} a series of reductions. Specifically, Schuch and Verstraete49 show that \( H_{\text{QMA}} \leq H_{\text{Heisenberg}} \leq H_{\text{Hubbard}} \leq H_{\text{elec}} \) where \( H_1 \leq H_2 \) means that instances of the ground state decision problem for Hamiltonian \( H_1 \) can be embedded (Karp reduced) into ground state problem instances of Hamiltonian \( H_2 \). In their proof, the authors utilize the magnetic field to encode the problem instances causing \( H_{\text{elec}} \) as defined in eqn (6) to include an additional local magnetic field, \( V_{\text{mag}} \). Note that the local field only affects the spin of the electron and does not require spin-dependent density functionals since the charge and spin do not couple. The Hamiltonian, \( H_{\text{QMA}} \), was listed earlier in eqn (9). Again since the reduction relies on the QMA-completeness of the 2-\textsc{local-spin Hamiltonian} problem, the promised error tolerance and the upper-bound of the coefficients of the Hamiltonian are required for QMA-completeness.

The \( H_{\text{Hubbard}} \) and \( H_{\text{Heisenberg}} \) Hamiltonians are commonly encountered models in condensed matter physics and are of
the form

\begin{equation}
H_{\text{Hubbard}} = \sum_{<ij>} \sum_{s \in \{\uparrow, \downarrow\}} t_{ij} a_i^s a_j^s + \sum_{<ij>} U_{ij} a_i^\uparrow a_i^\downarrow a_j^\downarrow a_j^\uparrow \quad (18)
\end{equation}

\begin{equation}
H_{\text{Heisenberg}} = \sum_{\sigma \in \{\uparrow, \downarrow\}} \sum_{<ij>} J_{ij} \sigma_i^\sigma_j^\ast \sigma_j^\sigma_i^\ast + \sum_{i} B^d_i \sigma_i^d \quad (19)
\end{equation}

The first Hamiltonian describes an electronic system where the Pauli matrices, \( \sigma^\dagger \), are expressed using orbital pairs as in eqn (10), while the second Hamiltonian describes a system of quantum spins. In both Hamiltonians, different problem instances are embedded using the local magnetic field. The embedding of \( H_{\text{Heisenberg}} \) instances into \( H_{\text{Hubbard}} \) follows directly from the orbital pair mapping described earlier in Section 3.3. The remaining two Karp reductions are more involved.

First, let us consider the reduction \( H_{\text{QMA}} \leq H_{\text{Heisenberg}} \). This Karp reduction follows along the same lines used in reducing \( k\text{QMA} \) from \( k = 5 \) to \( k = 2 \) based on perturbative gadgets widely used in quantum complexity proofs. A mediator spin splits the state space of the system into a low energy and a high energy sectors. In the low energy sector, the perturbative coupling to the high energy states “mediates” new interactions in the low energy sector. For example, with \( H_m = B_m | \phi_m \rangle \langle \phi_m | \) acting on the mediator spin and a perturbation \( V = \sigma_i^\dagger \sigma_m^\ast \sigma_m^\dagger \sigma_j^\dagger \), in the low energy space of spin \( m \), there is, at the second order correction, an effective interaction: \( \sigma_i^\dagger \sigma_j^\dagger \). Since the Hamiltonian problems refer to the ground state energy, the high energy sector is not important.

The reduction \( H_{\text{QMA}} \leq H_{\text{Heisenberg}} \) requires:

1. converting arbitrary strength couplings to constant strength couplings: \( J_{ij} \sigma_i^\dagger \sigma_j^\dagger - J \sigma_i^\ast \sigma_j^\ast \);
2. converting inequivalent couplings to equivalent couplings: \( \sigma_i^\dagger \sigma_j^\dagger - \sigma_i^\dagger \sigma_j^\dagger = \sigma_i^\dagger \sigma_i^\dagger + \sigma_j^\dagger \sigma_j^\dagger \); and finally converting equivalent couplings to Heisenberg interactions: \( \sigma_i^\dagger \sigma_j^\dagger - \sum_d \sigma_i^d \sigma_i^d + \sum_d \sigma_j^d \sigma_j^d \).

The full transformation requires 15 mediator spins where the local field on each of the mediator spins splits the system into low and high energy sectors. The specific local field depends on the interaction desired. The strength of the local field applied to the mediator spin \( B_m \) is usually very large to ensure that perturbation theory applies. So far this has limited the practical relevance of these constructions; for instance, field strengths required on the final set of mediator spins scales at nearly the 100th power of the system size.

The remaining reduction, \( H_{\text{Hubbard}} \leq H_{\text{elect}} \) was heuristically known since the Hubbard model phenomenologically describes electrons in solid state systems. Schuch and Verstraete rigorously demonstrate this reduction by careful accounting for error terms. They begin with a simple model used for studying solids where non-interacting (spin-less) electrons are subjected to a periodic delta function potential with \( M/2 \) such sites. The orbitals of this system can be solved exactly. The resulting second quantized Hamiltonian has uniform kinetic hopping terms: \( \sum_{<ij>} t_{ij} a_i^s a_j^s \). After incorporating the electron spin, electron-electron interactions are introduced. The strength of this interaction is rescaled by changing the spatial distance between neighboring sites until only the electrons at the same site can interact. Since each of the \( M/2 \) identical sites can only support one bound state, the exchange integral must vanish and the Coulomb integral is the same for each interaction yielding \( \sum_{i} U_{ij} a_i^\uparrow a_i^\downarrow a_j^\downarrow a_j^\uparrow \).

Since the Hubbard model and the Heisenberg model require magnetic fields to define problem instances, the electronic Hamiltonian which they consider is not precisely the same as eqn (6); this also requires that the functional takes into account the separate spin components of the probability density when performing the minimization in eqn (17).

Regardless, a polynomial time solution of \textsc{Universal Functional} would also solve the QMA-hard electronic Hamiltonian with local magnetic fields. This follows as the functional is convex

\begin{equation}
F \left[ \sum p_j \rho_j \right] = \min_{\rho^{(N)}} \text{Tr} \left[ (T_e + V_{ee}) \rho^{(N)} \right]
\end{equation}

and the one-electron probability densities are also convex allowing one to perform convex optimization to find the minimum energy of the QMA-hard electronic Hamiltonian with the local magnetic fields. Since the conditions for consistency are simple to check for one-electron probability densities, the complications encountered in 2-RDM convex optimization from the approximate consistency conditions are not present.

The inclusion of \textsc{Universal Functional} in complexity class QMA follows along the same lines of the generic two-fermion problem discussed in Section 3.4. However, in this case, Arthur must subtract the energy of the local field from the total energy to decide if the energy of \( F \) is above or below \( E_T \pm \delta \).

### 6.2 Restriction to pure states

The pure state restriction of DFT affects the complexity of computing \( F \). Consider evaluating \( F[\rho] \) where the input density arises from optimization over pure states \( |\Psi\rangle \) instead of mixed states as in eqn (17). In this case, the universal functional is no longer convex.

\[ \sum_{j} p_{j} \min_{\eta_{j}} \langle \Psi | (T_e + V_{ee}) | \Psi \rangle \neq \min_{\eta} \sum_{j} p_{j} \langle \Psi | (T_e + V_{ee}) | \Psi \rangle. \]

The optimization problem would then be NP and an oracle for \textsc{Universal Functional} would allow a Turing reduction from QMA to NP. Stated differently, an NP machine with access to an oracle for \textsc{Universal Functional}, could solve any problem in QMA. Similar to the 2-RDM case, the restriction to pure states complicates the verification such that two Merlins are required to show that the state is pure and that the proposed solution is correct.
7 Related methods and other results

In this article, we have focused on three pillars of quantum chemistry but there are other results and methods worth mentioning. First, we will discuss some consequences of the problems that have already been shown to be computationally difficult and then discuss methods that were thus far omitted.

In quantum chemistry, multi-determinant wave functions are useful for capturing static correlations which are important in many situations such as bond breaking. Common techniques used for such analysis are multi-configuration, complete active space, or restricted active space self-consistent field methods. As a consequence of Hartree–Fock being NP-complete, these methods are also at least NP-hard since they all include the single determinant Hartree–Fock calculation as a sub-case.

Perturbation theory also plays a prominent role in quantum chemistry when dynamical correlations need to be captured. At the forefront of these methods is Møller–Plesset (MP) perturbation theory and coupled cluster calculations. The MPn series is based on perturbing the Hartree–Fock solution using the difference between the exact Hamiltonian and the effective Fock matrix which includes the self-consistent mean-field. Including perturbative corrections thus requires that one could solve Hartee–Fock, but again the NP-completeness of these problems implies the NP-hardness of the MPn series. The same arguments hold for coupled-cluster calculations predicated on access to the Hartree–Fock state. In principle, the coupled-cluster expansion could start from any initial state so this argument does not fully suffice as the basis for a proof about the complexity of coupled-cluster techniques. It does indicate that, at the very least, the standard approaches would be limited by the Hartree–Fock computation in the first place.

Now we turn to the discussion of other notable methods omitted earlier. First, Troyer and Wiese have shown that the sign problem of quantum Monte Carlo, when properly defined, is NP-hard. They showed that the stochastic evaluation of the sign problem of quantum Monte Carlo, when properly defined, can only be done to polynomial accuracy. Including perturbative corrections thus requires that one could solve Hartree–Fock, but again the NP-completeness of these problems implies the NP-hardness of the MPn series. The same arguments hold for coupled-cluster calculations predicated on access to the Hartree–Fock state. In principle, the coupled-cluster expansion could start from any initial state so this argument does not fully suffice as the basis for a proof about the complexity of coupled-cluster techniques. It does indicate that, at the very least, the standard approaches would be limited by the Hartree–Fock computation in the first place.

With respect to complexity in quantum chemistry, we have examined, in detail, the computational complexity of three different quantum chemistry algorithms: Hartree–Fock, N-Rep, and Universal Functional. The difficulty of these computational chemistry problems relative to problems found in the same complexity class as computing the classical partition function. The complexity class of these problems is #P where the problem instances request the number of solutions to problem in complexity class P.

8 Remarks on approximability of quantum systems

Before closing the paper, we make a few remarks on scaling of the error tolerances. In the computational problems investigated, the error tolerance was upper bounded by an inverse polynomial $1/q(N)$ in the system size. In Hartree–Fock, it was the pseudo-spin mapping that necessitated the promise, but, in Universal Functional and N-Rep, the promise is inherited from the 2-LOCAL-SPIN HAMILTONIAN problem. At first glance, it would appear that the error tolerance becomes more restrictive as the system size increases, however this is not necessarily the case.

We focus on the problem $k$-LOCAL-SPIN HAMILTONIAN since it is at the heart of the reductions for both Universal Functional and N-Rep. In this problem, the error tolerance, $\delta$, is normalized by the number of terms in the Hamiltonian,$^{75,76} m$. Therefore, the error tolerance only shrinks with system size if the number of non-zero terms in the Hamiltonian is constant or grows slower than $q(N)$. In many cases, the number of terms in the Hamiltonian increase with system size. Thus, even if asking for fixed error, e.g. 1 kcal mol$^{-1}$, as the system size increases the promise is still fulfilled and the problem of deciding instances of $k$-LOCAL-SPIN HAMILTONIAN is QMA-complete.

When the error tolerance is independent of the system size or grows slowly, it is not clear if the problem remains QMA-complete. This is related to the on-going research into possible quantum generalizations of the PCP theorem.$^{88}$ The PCP theorem shows that the task of approximating some NP-complete problems is also NP-complete.$^{78,79}$ A quantum generalization, if it exists, would allow one to show that $k$-LOCAL-SPIN HAMILTONIAN remains QMA-complete even when the normalized error tolerance is bounded by a constant instead of an inverse polynomial of $N$. This remains a prominent open question in modern computer science and is discussed more thoroughly by Osborne.$^{19}$

9 Outlook

With respect to complexity in quantum chemistry, we have examined, in detail, the computational complexity of three different quantum chemistry algorithms: Hartree–Fock, N-Rep, and Universal Functional. The difficulty of these computational chemistry problems relative to problems found in

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$\dagger$ One may ask why not use the norm of the Hamiltonian? For a Hamiltonian, $H$, the operator norm is the maximum eigenvalue and computing the maximum eigenvalue is just as hard as computing the ground state of $-H$, thus also a difficult problem.

$\S$ The term PCP refers to probabilistically checkable proof systems where the verifier is only allowed to randomly access parts of a non-deterministically generated proof. Using only polynomial time and a fixed amount of random bits, the verifier is to validate or invalidate the proof.
other fields gives insights into why computational chemistry is difficult. The difficulty of these problems imply that even a quantum computer will be unable to solve all instances since it is believed that BQP does not contain NP. However, even if quantum computers are never constructed, the study of quantum computational complexity gives new insight into the relative difficulty of problems. For instance, characterizing valid two-electron reduced density matrices or evaluating the universal functional of DFT is arguably more difficult than evaluating the Hartree–Fock energy based on the probable separation of complexity classes QMA and NP.

This article aims to facilitate cross-fertilization of ideas between numerical chemical physics and quantum computational complexity by providing an interdisciplinary perspective. While the picture is far from complete, it does provide a new purview of computational quantum chemistry. For developers and users of software packages, these results should not change their ambitions, only challenge them. Just as advances in numerical simulations have assisted experimental chemistry by informing, interpreting and complementing physical data, we expect in the future, an expanded understanding of computational complexity can provide added value to numerical data from computational simulations.

Despite the new results presented, there are many remaining questions essential to the expansion of applications of theoretical computer science to chemistry.

- **What is the computational complexity of other molecular properties besides energy?** The computational difficulty of properties such as excited states or derivatives of the energy have not been systematically investigated yet. The examination of other properties of molecular systems will provide fertile ground for future research into the difficulty of quantum chemistry.

- **Can studying open system dynamics simplify the complexity?** That is, if one is only interested in the electronic properties of a fragment of a system, e.g. a solute molecule or a molecule in heat or phonon bath, does the computational complexity change?

- **Although problems (i.e. Hartree–Fock) are difficult in the worst case, what precise restrictions or well defined classes of molecules admit tractable computations?** The answer to this questions is known intuitively by many researchers who have performed thousands of electronic structure calculations. Is the restriction to the first three-rows of the periodic table enough to ensure all calculations converge reasonably quickly? What are the most general restrictions that allow such a promise to be fulfilled?

- **Do QMA-hard scalar electronic potentials exist?** In other words, what is the computational complexity of Coulombic Hamiltonians without magnetic fields? The proof by Schuch and Verstraete, encodes problem instances into magnetic Hamiltonians without magnetic fields? The proof by Schuch

References
