Simulating Quantum Mechanics on a Quantum Computer*

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Abstract

Algorithms are described for efficiently simulating quantum mechanical systems on quantum computers. A class of algorithms for simulating the Schrödinger equation for interacting many-body systems are presented in some detail. These algorithms would make it possible to simulate nonrelativistic quantum systems on a quantum computer with an exponential speedup compared to simulations on classical computers. Issues involved in simulating relativistic systems of Dirac and gauge particles are discussed.

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1 Introduction

There has been a great deal of excitement in the field of quantum computing over the last few years, due largely to the result of Shor [1] showing that large numbers can be factored on a quantum computer in a time that scales polynomially with the number of digits. Shor’s work was based on earlier results showing that certain oracle problems could be solved on a quantum computer exponentially faster than on a classical computer [2]. Since it is believed that factoring is an exponentially hard problem on a classical computer, Shor’s result provided for the first time a concrete demonstration that quantum computers can achieve useful results much more efficiently than classical computers. Despite a great deal of work in this field, however, there has been little progress toward discovering other algorithms which give the quantum computer an exponential advantage in performance.

Early in the development of quantum computing, it was suggested by Feynman [3] that it might be possible to simulate quantum mechanical systems exponentially faster on quantum computers than on classical computers. Feynman argued that understanding this issue might help to quantify the differences between the computational capacities of classical and quantum computers. Recently, it was argued by Lloyd [4] that for a fairly general class of quantum systems, particularly for discrete systems, it should be possible to achieve an exponential speedup when simulating quantum mechanical systems on a quantum computer. In this paper we will discuss algorithms which are concrete realizations of these general arguments.

There are three particular types of quantum theories which we would like to try to simulate on a quantum computer:

- Nonrelativistic many-body systems (the Schrödinger equation)
- Relativistic many-fermion systems (the many-body Dirac equation)
- Gauge field theories (quantum Yang-Mills theories)

In this paper we describe in some detail a class of algorithms for simulating the many-body Schrödinger equation. These algorithms were presented from a different point of view in
Based on simple Quantum Cellular Automata (QCA) [7] and Quantum Lattice-Gas Automata (QLGA) [8] models, these algorithms can be used to simulate systems of interacting nonrelativistic quantum particles with speedup exponential in the number of particles in the system. We give a short discussion of progress which has been made toward developing algorithms for simulating systems of many Dirac particles, and we discuss prospects for simulating gauge field theories on a quantum computer.

2 Quantum computers

We begin with a brief description of a universal quantum computer, in order to fix notation. For recent reviews of quantum computing in general, see [9]. The state space of a quantum computer is defined to be the Hilbert space associated with a finite number ($N$) of two-state quantum bits ($q$-bits). A natural basis for this Hilbert space is given by the set of $2^N$ states

$$\left| \sigma \right> = \left| \sigma_1 \sigma_2 \ldots \sigma_N \right>$$

where $\sigma_i \in \{\uparrow, \downarrow\}$.

A single unit of computation is defined to be the action of an arbitrary two-bit gate on the state of the system. Mathematically, this corresponds to acting on the state of the system with a $2^N \times 2^N$ matrix which is a tensor product of an arbitrary $4 \times 4$ unitary matrix (acting on the Hilbert space associated with a pair of $q$-bits $\sigma_i, \sigma_j$), with an identity operator of dimension $2^N - 2 \times 2^N - 2$. It has been shown [10] that such two-bit gates are universal for quantum computation.

A measurement in a quantum computer is performed by measuring the states of some of the $q$-bits. It is sufficient to restrict attention to measurements of the states of individual spins with respect to the canonical basis. As dictated by quantum mechanics, a single $q$-bit in the quantum state $\left( \psi_{\uparrow} | \uparrow \rangle + \psi_{\downarrow} | \downarrow \rangle \right)$ will be measured to be in the state $\uparrow$ (or $\downarrow$) with probability $|\psi_{\uparrow}|^2$ (or $|\psi_{\downarrow}|^2$).

The definition of a universal quantum computer is closely modeled on the conceptual framework of classical computation. It has been suggested [11] that a more general definition of a quantum computer may be desirable, as there may be physical systems which exhibit the
capacity for performing useful quantum computation which do not fit into this axiomatic framework. We will discuss briefly in the following section an example of a computation which is difficult on a standard universal quantum computer, which might be more easily implemented on a generalized quantum computer. However, we will otherwise remain within the context of the standard definition of a universal quantum computer in terms of q-bits and two q-bit operations.

3 Simulating the Schrödinger equation

In this section we describe a class of algorithms for simulating the many-body Schrödinger equation on a quantum computer with exponential speedup over simulations on a classical computer. The algorithms we discuss here are all based on a second-quantized formalism which naturally fits into the framework of quantum field theory. Another approach to simulating the many-body Schrödinger equation was discussed in [12].

To begin the discussion, let us recall the form of the Schrödinger equation for a system of interacting particles moving in $d$ dimensions.

3.1 Review of Schrödinger equation

The Schrödinger equation for a single free particle of mass $m$ moving in $d$ dimensions is

$$i \frac{\partial}{\partial t} \psi(x, t) = -\frac{1}{2m} \sum_i \frac{\partial^2}{\partial (x^i)^2} \psi(x, t)$$

(2)

where we have chosen units with the Planck constant set to be $\hbar = 1$. If the particle is moving in the presence of an external potential $U(x)$, the Schrödinger equation becomes

$$i \frac{\partial}{\partial t} \psi(x, t) = -\frac{1}{2m} \sum_i \frac{\partial^2}{\partial (x^i)^2} \psi(x, t) + U(x) \psi(x, t).$$

(3)

For a system of $n$ identical particles interacting via a (symmetric) pairwise potential $U(x, x_j)$, the Schrödinger equation is

$$i \frac{\partial}{\partial t} \psi(x_1, \ldots, x_n, t) = \sum_{k=1}^n -\frac{1}{2m} \sum_{i=1}^d \frac{\partial^2}{\partial (x^i_k)^2} \psi(x_1, \ldots, x_n, t)$$

(4)
$$+ \sum_{j<k} U(x_j, x_k) \psi(x_1, \ldots, x_n, t)$$

If we wish to simulate the $n$-particle Schrödinger equation in $d$ dimensions, one approach is to discretize space. If we discretize so that the particles move on a spatial lattice with $l$ lattice sites in each direction, the number of independent components of the $n$-particle wavefunction grows as $l^{dn}$. Even for $d = 3$, for reasonably large values of $l$ and $n$ this number becomes extremely large. If we wish to simulate the system on a classical computer, the number of independent components in the wavefunction is a lower bound both on the amount of memory needed to store the state of the system at a fixed point in time, and also for the amount of computation needed to take a single time step in the simulation. For a lattice size of $l = 20$ and a system of 20 particles, we have $l^{3n} = 20^{60} \sim 10^{78}$, which is clearly far beyond the memory and computational resources of any imaginable classical computer. In the following sections we will describe algorithms with which this system can be simulated on a quantum computer with memory and computational requirements per time step on the order of $l^d$, independent of $n$ (so long as $n \ll l^d$).

### 3.2 Simulating a free Schrödinger particle in one dimension

We begin our discussion of simulations on a quantum computer by considering the simplest case: a free Schrödinger particle moving on a lattice in one dimension. Let us consider a one-dimensional lattice with $l$ vertices. We will associate a single q-bit $\sigma_i$ with each of the vertices $i$. Let us associate a q-bit $\sigma_i$ in the state $\uparrow$ with the presence of a “particle” at the $i$th lattice site, and a q-bit in the state $\downarrow$ with the absence of a particle. The canonical basis for the Hilbert space contains $2^l$ states, with anywhere between 0 and $l$ particles in the available states. Let us restrict attention to the single-particle subspace of the Hilbert space. This subspace has a basis containing $l$ states; we denote by $|k\rangle$ the state where the single particle is contained at lattice site $k$. Graphically, this state would look like

$$\cdots \downarrow \downarrow \downarrow \uparrow \downarrow \downarrow \downarrow \cdots$$

$$\cdots \ k-1 \ k \ k+1 \ \cdots$$
We now wish to define a discrete time dynamics on the Hilbert space which gives a Schrödinger equation in the continuum limit. Writing the wave function describing the single-particle state at time \( t \) in the form

\[
\psi(t) = \sum_i \psi_i(t) |i\rangle,
\]

(5)

the equation of motion for evolution over a single time step will be

\[
\psi(t + 1) = M \cdot \psi(t)
\]

(6)

where \( M \) is a unitary matrix. One way of getting a Schrödinger equation would be to find an operation on the full Hilbert space which had the effect in the single particle Hilbert space of acting by the operator

\[
M = \begin{pmatrix}
    b & a & 0 & 0 & \cdots & 0 & 0 & a \\
    a & b & a & 0 & \cdots & 0 & 0 & 0 \\
    0 & a & b & a & \cdots & 0 & 0 & 0 \\
    0 & 0 & a & b & \cdots & 0 & 0 & 0 \\
    \vdots & \ddots & \ddots & \ddots & \ddots & \ddots & \ddots & \ddots \\
    0 & 0 & 0 & \cdots & b & a & 0 \\
    0 & 0 & 0 & \cdots & a & b & a \\
    a & 0 & 0 & \cdots & 0 & 0 & a & b
\end{pmatrix}
\]

(7)

where \( a, b \) are complex numbers satisfying \(|b|^2 + 2|a|^2| = 1\) and \( a\bar{b} + \bar{a}b = 0 \). Unfortunately, this operator cannot be implemented by a simple set of two q-bit operations [7, 8]. One way of seeing that this is not possible is to take the inverse of the matrix \( M \). The matrix \( M^{-1} \) is a dense matrix for generic lattice size \( l \), indicating that \( M \) cannot be implemented by performing some sequence of local two q-bit operations at each lattice site in a way which is independent of \( l \).

It is worth noting at this point that although the operation \( M \) cannot be simply realized in the framework of universal quantum computers using q-bits and two q-bit operations, there are simple physical systems which behave in a fashion very similar to the dynamics which
this operation would define. In particular, a lattice of spin-1/2 particles with a Hamiltonian proportional to the sum of exchange operations, as discussed in [13], would lead to a very similar dynamics. Thus, it may be advantageous to consider a wider range of quantum computational devices in searching for algorithms for simulating generic quantum systems.

Returning to the framework of universal quantum computers, we now define a slightly more complicated dynamics which can be implemented using two q-bit operations, and which leads to the desired Schrödinger equation. Consider the two operators acting on the single particle Hilbert space

\[
M_1 = \begin{pmatrix}
  b & a & 0 & 0 & \cdots & 0 & 0 & 0 \\
  a & b & 0 & 0 & \cdots & 0 & 0 & 0 \\
  0 & 0 & b & a & \cdots & 0 & 0 & 0 \\
  0 & 0 & a & b & \cdots & 0 & 0 & 0 \\
  \vdots & \ddots & \vdots & & \ddots & \vdots & \vdots & \vdots \\
  0 & 0 & 0 & 0 & \cdots & b & a & 0 \\
  0 & 0 & 0 & 0 & \cdots & a & b & 0 \\
  0 & 0 & 0 & 0 & \cdots & 0 & 0 & b \\
  0 & 0 & 0 & 0 & \cdots & 0 & 0 & a \\
\end{pmatrix}
\]

\[
M_2 = \begin{pmatrix}
  b & 0 & 0 & 0 & \cdots & 0 & 0 & 0 & a \\
  0 & b & a & 0 & \cdots & 0 & 0 & 0 & 0 \\
  0 & a & b & 0 & \cdots & 0 & 0 & 0 & 0 \\
  0 & 0 & 0 & b & \cdots & 0 & 0 & 0 & 0 \\
  \vdots & \ddots & \vdots & & \ddots & \vdots & \vdots & \vdots & \vdots \\
  0 & 0 & 0 & 0 & \cdots & b & 0 & 0 & 0 \\
  0 & 0 & 0 & 0 & \cdots & 0 & b & a & 0 \\
  0 & 0 & 0 & 0 & \cdots & 0 & a & b & 0 \\
  a & 0 & 0 & 0 & \cdots & 0 & 0 & 0 & b \\
\end{pmatrix}
\]

(8)

where \(|a|^2 + |b|^2 = 1\) and \(ab + ba = 0\). We can now describe a quantum algorithm which gives a time-evolution equation for the single-particle sector

\[
\psi(t) = M_1 \cdot M_2 \cdot \psi(t - 2)
\]

(9)

The essential point is that the single-particle state can be transformed by either of the matrices \(M_i\) by acting on the particles in pairs by the two q-bit operator which is described in the basis \(\{|↓↓\rangle, |↑↓\rangle, |↓↑\rangle, |↑↑\rangle\}\) by the matrix

\[
s = \begin{pmatrix}
  1 & 0 & 0 & 0 \\
  0 & b & a & 0 \\
  0 & a & b & 0 \\
  0 & 0 & 0 & 1 \\
\end{pmatrix}
\]

(10)
Graphically, we can describe the implementation of the dynamics (9) through the following sequence of applications of the operator \( s \) to pairs of q-bits

\[
\begin{array}{cccc}
t & \quad & t + 1 & \quad & t + 2 & \quad & t + 3 \\
\begin{array}{cccc}
s & \quad & s & \quad & s & \quad & s \\
S & \quad & S & \quad & S & \quad & S \\
s & \quad & s & \quad & s & \quad & s \\
S & \quad & S & \quad & S & \quad & S \\
\end{array}
\end{array}
\]

We will now take the continuum limit of the dynamics (9) as the lattice spacing \( \epsilon \) becomes small. In the continuum limit, we are interested in the behavior of wavefunctions which are continuous and which therefore vary only by order \( \epsilon \) between adjacent lattice sites. We can analyze the dynamics of these wavefunctions in the continuum limit by performing a mode analysis. Assume that at time \( t = 0 \), we act with the matrix \( M_2 \) which acts on the pairs of q-bits \((2j, 2j+1)\). If at time \( t = 0 \) the even and odd lattice sites respectively have amplitudes given by

\[
\begin{align*}
\psi_{2j}(0) &= \alpha e^{i\kappa(2j)\epsilon} \\
\psi_{2j+1}(0) &= \beta e^{i\kappa(2j+1)\epsilon}
\end{align*}
\]

Then at time \( t = 2 \) we have

\[
\begin{pmatrix}
\psi_{2j}(2) \\
\psi_{2j+1}(2)
\end{pmatrix} =
\begin{pmatrix}
b^2 + a^2 e^{-2i\kappa\epsilon} & ab(e^{i\kappa\epsilon} + e^{-i\kappa\epsilon}) \\
ab(e^{i\kappa\epsilon} + e^{-i\kappa\epsilon}) & b^2 + a^2 e^{2i\kappa\epsilon}
\end{pmatrix}
\begin{pmatrix}
\psi_{2j}(0) \\
\psi_{2j+1}(0)
\end{pmatrix}.
\]

For a fixed value of the angular wave number \( \kappa \) the matrix which appears in this equation has two eigenvalues. In the continuum limit, where \( \kappa\epsilon \to 0 \), the eigenvectors of this matrix approach the limiting values \((1, 1)\) and \((1, -1)\). Since we are interested in the behavior of wavefunctions which are smooth to order \( \epsilon \) at \( t = 0 \), only the first of these eigenvectors will be relevant for describing time-development in the continuum limit. The eigenvalue associated with the first of these eigenvectors is

\[
\lambda(\kappa\epsilon) = b^2 + a^2 \cos(2\kappa\epsilon) + a \cos(\kappa\epsilon) \sqrt{4b^2 + 2a^2(\cos(2\kappa\epsilon) - 1)}
\]
Performing a power series expansion for small $\epsilon$ and fixed $\kappa$ we have

$$\lambda(\kappa\epsilon) = (a + b)^2(1 - \frac{a}{b}\kappa^2\epsilon^2 + \mathcal{O}(\epsilon^4))$$

(14)

By factoring out an overall time-dependent phase from the wavefunction

$$\Psi(x, t) = (a + b)^{-1}\psi(x, t)$$

(15)

we see that after scaling the time variable as $\epsilon^2$, the wavefunction $\Psi$ obeys the dispersion relation

$$i\omega = \frac{a}{2b}\kappa^2.$$  

(16)

Thus, in the continuum limit, to within an error of order $\epsilon$, the wavefunction $\Psi(x, t)$ obeys the Schrödinger equation

$$i\partial_t \Psi(x, t) = -\frac{1}{2m}\partial_x^2 \Psi(x, t)$$

(17)

for a particle of mass

$$m = \frac{bi}{a}.$$  

(18)

Note that this mass is always real, due to the condition $a\bar{b} + \bar{a}b = 0$ necessary for unitarity.

We have now seen that the dynamics (9) gives rise in the continuum limit to a Schrödinger equation for a single free particle moving in one dimension. A number of similar models have been considered previously by other authors. Essentially the same model was studied by Succi and Benzi [14]; a number of numerical experiments were performed on this model by Succi to study the emergent Schrödinger behavior in the continuum limit [15]. A closely related model was described by Feynman [16] in the context of the single particle Dirac equation in one spatial dimension. In Feynman’s model, however, the amplitude $b$ was taken to scale as the lattice spacing $\epsilon$. This model was the basis for the recent work of Meyer [8, 17, 18] on the many-particle one-dimensional Dirac system; we will discuss this system in Section 4.
3.3 External potential

We now have an algorithm for simulating a free quantum particle in one dimension. The dynamics of this system, which are described by the dispersion relation (16), are essentially trivial. We shall now add various features to the model to give a system of more physical interest. First, we describe how an external potential can be easily incorporated into the system.

Given an external potential function $U(x)$, the effect of this potential can be brought into the dynamics by acting on each q-bit at each time step with the matrix

$$U_x = \begin{pmatrix} 1 & 0 \\ 0 & e^{-ic^2U(x)} \end{pmatrix}$$

where $x$ is the position of the q-bit. The inclusion of this operation changes the time development of the system so that the differential equation satisfied by $\Psi$ becomes

$$i \partial_t \Psi(x,t) = -\frac{1}{2m} \partial_x^2 \Psi(x,t) + U(x)\Psi(x,t)$$

in the continuum limit, precisely the Schrödinger equation for a particle in an external potential.

It is worth noting that the continuum limit of the theory with an external potential may not be well-behaved when the external potential is not smooth. If the potential is singular or has a discontinuity, it is possible that the eigenvectors of the matrix in (12) with limiting form $(1, -1)$ can be excited. This issue is discussed by Meyer in [17]. As long as the potential function is smooth, however, the only eigenstates of the system which are excited will be of the $(1, 1)$ form, and the Schrödinger equation in an external potential will be satisfied to order $\epsilon$. We have performed numerical analyses of the spectrum for simple systems and have verified that for smooth potentials the behavior of the model closely matches the continuum system being modeled [6].
3.4 Multiple particles

Thus far, we have really not used the power of the quantum computer at all. The number of states available in the single-particle sector of the Hilbert space is essentially the same as the number of q-bits, so the algorithm for a single particle could just as well be implemented on a classical computer. Now, however, let us consider the behavior of the multiple-particle sectors of the Hilbert space with the same algorithm described above. Keeping the notation that a q-bit in the state $\uparrow$ corresponds to the presence of a particle, we can for example graphically describe a state with three particles at positions $k - 4, k, k + 2$ by

```
... ↓ ↓ ⋄ ↓ ↓ ⋄ ↓ ↓ ⋄ ↓ ↓ ⋄...
... k - 1 k k + 1 ...
```

Because the operator $s$ in (10) preserves particle number, the number of particles in the system is invariant under the dynamics. If we have a lattice of size $l$, as long as the number of particles present $n$ is small compared to the lattice size ($n \ll l$), generically particles will propagate for a fairly large number of time steps without encountering other particles. In the limit where the lattice spacing gets arbitrarily small but the number of particles remains fixed, we will obtain a Schrödinger equation for $n$ particles, which will be noninteracting except when a pair of particles are at the same lattice site. Thus, we have described an algorithm for simulating many Schrödinger particles in one dimension interacting with a local ($\delta$ function) interaction potential.

By incorporating an external potential using single-particle operators like (19) we can simulate a system of $n$ Schrödinger particles moving in an external potential. It is perhaps more interesting, however, to include interactions between the particles through a pairwise potential function $U(x, y)$. A local interaction corresponding to a potential $U(x, y) \sim \delta(x - y)$ with an arbitrary phase can be incorporated by including a phase $\phi$ in the two-particle component of the matrix $s$ in (10). In general, however, we would like to incorporate nonlocal potential functions. In order to implement the effects of a general interparticle potential, we can act at every time step on the four-dimensional Hilbert space associated with every pair
of q-bits at positions $x$ and $y$ with the operator

$$U_{x,y} = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & e^{-i \epsilon 2 U(x,y)}
\end{pmatrix}.$$ (21)

This operator acts on the state $\uparrow\uparrow$ with a nontrivial phase, and therefore affects only the basis states where there are particles at positions $x$ and $y$. Just as in the case of the external potential, incorporating the effects of these operators turns the dynamical equation for $\Psi$ into a Schrödinger equation for many interacting particles (4).

### 3.5 Systems in arbitrary dimension $d$

So far we have considered systems of particles moving only in a single spatial dimension. Of course, most problems of physical interest involve particles moving in three (or more) spatial directions. We will now discuss the generalization of the algorithms discussed above to higher dimensions. One simple way of simulating a $d$-dimensional system is to put the q-bits on a $d$-dimensional lattice, and to implement the kinetic part of the time-development rule by repeatedly using the algorithm described in section 3.2 along each of the dimensions successively. This leads to a fairly simple set of algorithms for simulating the many-particle interacting Schrödinger equation in $d$-dimensions. There are several aspects of this algorithm, however, which may be somewhat problematic. For one thing, acting in each of the directions in turn with the kinetic operator (9) breaks the underlying lattice symmetry. For another thing, the number of q-bits which behave differently under the time-development rule scales as $2^d$, which for large $d$ may complicate the numerical interpretation of simulations.

A nice way of packaging the q-bits and kinetic time-development rule in $d$-dimensions is given by the general class of systems called Quantum Lattice-Gas Automata (QLGA) [8, 5]. These models are defined by analogy with classical lattice-gas automata [19] to be simple dynamical systems in which q-bits on a lattice undergo a time-development which consists of alternating propagation and collision steps. At each lattice site there are a set of q-bits
Figure 1: A typical basis state in the Hilbert space of a 2D QLGA. Filled circles correspond to occupied particle states (q-bits in the state $\uparrow$). Arrows correspond to directions of propagation.

corresponding to particle occupation sites for particles with velocity vectors lying on the lattice. On a cartesian lattice in $d$-dimensions, for example, we could have $2d$ q-bits at each lattice site, corresponding to sites for particles moving along each of the basis vectors. Just as in the previous models, we associate a q-bit in state $\uparrow$ ($\downarrow$) with the presence (absence) of a particle. These models have the advantage that they can be defined in a fashion which is completely symmetric with respect to the discrete symmetry group of the underlying lattice. Also, the number of distinct types of q-bits goes as $2^d$ instead of $2^{2d}$ as in the iterative model mentioned above. An example of a basis state for the Hilbert space of a QLGA in two dimensions is given in Figure 1. The filled circles represent occupied particle sites, and the empty circles represent unoccupied sites.

In the propagation stage of the time-development of a QLGA, the state of the system is transformed by a permutation on the q-bits. This permutation takes the q-bit associated with each position and velocity to a q-bit at the lattice site reached by adding the velocity vector to the position. As indicated by the arrows in the example in Figure 1, this transformation can be achieved by exchanging the q-bit states in pairs. For each pair of q-bits which must be exchanged, the operator acting on the Hilbert space is given in the basis $\{|\downarrow\downarrow\rangle, |\uparrow\downarrow\rangle, |\downarrow\uparrow\rangle\}$. 
by the matrix
\[
E = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & 0 & 1 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 0 & 1
\end{pmatrix}
\] (22)

In the collision stage of the time-development, the state associated with all the q-bits at each lattice site is acted on by a unitary “collision” operator. This operator is chosen to conserve particle number. In general, we would like to choose a collision operator which is invariant under the group of discrete symmetries of the lattice. As in the one-dimensional model discussed above, for systems where the particle number is fixed and the lattice spacing becomes small, the part of the collision operator in the single-particle sector determines the propagation of the individual particles. On a cartesian lattice in \(d\) dimensions, the collision operator in the single particle sector is parameterized by 3 complex phases \(\mu, \nu, \lambda\). These phases correspond to the eigenvalues of the collision matrix for vectors which lie in the three irreducible representations of the discrete rotation group into which the \(2d\)-dimensional single particle space decomposes. Specifically, \(\mu\) is the eigenvalue associated with the constant vector \((1, 1, \ldots, 1)\) and \(\nu\) is the eigenvalue of vectors which are antisymmetric under a parity transformation. It is shown in [5] that in the generic case where the three eigenvalues are distinct, the behavior of the individual particles in the system is governed by a Schrödinger equation for a nonrelativistic particle with mass \(m\) given by the relation
\[
\frac{i}{2m} = \frac{1}{d} \left(\frac{\nu}{\mu - \nu} + \frac{1}{2}\right).
\] (23)

By combining the propagation and collision processes into a single time-development rule, we have a QLGA model which has a continuum limit described by a many-particle Schrödinger equation in \(d\) dimensions. As described so far, the particles in this system interact only through local (\(\delta\) function) interactions, which are parameterized by the part of the collision matrix acting in the multiple-particle space. Just as in the one-dimensional model discussed above, it is straightforward to include external and interparticle potentials,
so that an arbitrary system of nonrelativistic interacting quantum particles can be simulated using this type of model.

In fact, the one-dimensional model discussed in sections 3.2-3.4 is precisely equivalent to a simple one-dimensional QLGA. We can associate the even and odd lattice sites with right- and left-moving particle sites at even time steps, and with left- and right-moving particle sites at odd time steps. If we define a one-dimensional QLGA with a collision matrix given by $s$ as in (10), it is easy to see that the QLGA decomposes into two noninteracting copies of the one-dimensional model defined through (9). For this choice of collision matrix we have $\mu = a + b$ and $\nu = a - b$ (the sign on $\nu$ arises because the collision operator $s$ implicitly changes the direction of velocity for each particle). Inserting these values for $\mu, \nu$ into (23) reproduces (18) as we would expect.

It should be noted that the algorithms we have discussed here are most easily used to simulate systems of interacting bosons. The microscopic quantum system only contains a single state in the Hilbert space corresponding to each configuration of particles. Two paths in which particles move to positions $x, y$ and $y, x$ will contribute amplitudes which add with no change of sign, as is appropriate for Bose statistics. It is possible to modify the algorithm to incorporate Fermi statistics, however this requires somewhat more bookkeeping. This issue was discussed in [20].

### 3.6 Computational complexity

We have now described a class of algorithms for simulating the nonrelativistic many-body Schrödinger equation on a quantum computer. As discussed in Section 3.1, simulating $n$ quantum particles on a lattice of size $l^d$ requires a computation of complexity $O(l^{dn})$ on a classical computer. We now briefly discuss the time and memory requirements for such a simulation on a quantum computer. Using the quantum lattice-gas automaton formalism, we can simulate such a system with $2d \cdot l^d$ q-bits. The propagation step takes on the order of $dl^d$ two q-bit operations, since the q-bits are simply exchanged pairwise. The free kinetic collision step requires on the order of $d^2l^d$ operations (the factor of $d^2$ is needed to construct
the collision operator at each vertex from two q-bit operations). Incorporating an arbitrary interparticle interaction potential would require on the order of $d^2 l^2 d$ operations, since we must act on each pair of q-bits independently. This operation will clearly be the most computationally expensive part of the simulation. We see, however, that the memory and time requirements for the algorithms we have discussed are only polynomial in the lattice size, and are completely independent of the number of particles as long as $n \ll l^d$. Thus, we can achieve a speedup for many-particle quantum simulations which is exponential in the number of particles being simulated. For example, for a lattice of size $l = 20$ in three spatial dimensions, the number of operations needed in a simulation would be on the order of $36 \cdot 2^{10} \sim 2.3 \cdot 10^9$ compared with $10^{78}$ operations for a simulation on a classical computer.

In discussing the computational complexity of these simulation algorithms, it is worth noting that if we simulate nonrelativistic fermions extra work must be done to keep track of the phase of the state. Using the bookkeeping method suggested in [20], for example, we must define a canonical ordering for all the particles, and we must check to see when the propagation changes the ordering of any pair of particles. This will take roughly on the order of $d^2 l^2 d$ operations per time step, just as for the interparticle interaction. In fact, however, we can incorporate this phase into the interparticle interaction matrix, so that the simulation of an interacting system of nonrelativistic fermions can be achieved in precisely the same number of quantum operations as the simulation of interacting “hard” bosons (particles with bosonic statistics which cannot occupy the same lattice site).

4 Simulating relativistic systems

In the bulk of this paper we have discussed the simulation of nonrelativistic systems of many Schrödinger particles. Many systems of physical interest are described to a high degree of precision by the nonrelativistic many-body Schrödinger equation. However, there are many physical systems for which the Schrödinger equation is not a sufficiently accurate model. In particular, it is desirable to have a way of simulating relativistic quantum systems such as
gauge fields or Dirac fermions.

Let us first consider the problem of simulating free Dirac fermions in \( d \) dimensions on a quantum computer. As mentioned in Section 3.2, it was found by Feynman [16] that if the parameter \( b \) is taken to approach 0 at the same rate as the lattice spacing, the one-dimensional model described above satisfies the first-order two-component Dirac equation. By generalizing this single-particle model to the QLGA framework, it was shown by Meyer [8] that a free system of many noninteracting Dirac particles in \( 1 + 1 \) dimensions could be simulated on a quantum computer.

The problem of simulating Dirac fermions on a lattice in more than one spatial dimension has presented difficulties to physicists for decades now. A full discussion of the approaches which have been used to try to solve this problem is beyond the scope of this paper. The best-known obstacle to a sensible discretization of a field theory of Dirac particles is the fermion doubling problem. If we try naively to generalize Feynman’s model for the one-dimensional Dirac system to higher dimensions, we immediately encounter several difficulties. First, even for a single particle it is not straightforward to find a lattice rule which gives the propagator for the Dirac equation. Within the QLGA framework, in fact, it is not possible to construct a single-particle Dirac equation in any dimension \( d > 1 \). A model suggested several years ago by Bialynicki-Birula [21], however, fits very closely into the framework we have been discussing, and may provide a way of simulating systems of many Dirac particles without encountering the difficulties which have plagued previous efforts.

The basic approach of Bialynicki-Birula is to construct a quantum cellular automaton model with a single time-development rule which depends only on local variables. He finds that in three spatial dimensions, on the body-centered cubic lattice where every vertex has eight neighbors, it is possible to find such a rule whose continuous limit corresponds to a two-component Weyl spinor. Remarkably, it seems that such a model cannot exist on the three-dimensional cartesian lattice. A Dirac particle is formed by taking two Weyl spinors and including a mass term by hand. As described by Bialynicki-Birula, the model avoids the no-go theorem of Nielsen and Ninomiya [22] because it has discrete time evolution.
The model of Bialynicki-Birula suffers from the same difficulty as the model with time-development operator (7) which we discussed in Section 3.2, and thus this algorithm cannot be implemented naturally on a universal quantum computer. However, the existence of the model suggests that by using a multi-step rule on the body-centered cubic lattice, it might be possible to efficiently simulate free Dirac fermions in three dimensions on a quantum computer.

If one could indeed simulate many free Dirac particles on the lattice, it is natural to try to simulate a full second-quantized field theory of Dirac particles. To do this, one would need to introduce antiparticles corresponding to negative energy states. Incorporating a potential directly into a Dirac theory is liable to be problematic, however, since the simplest interaction term, the four-fermi interaction, is nonrenormalizable and would destroy the continuum limit of the theory [23].

Of course, if one wishes to simulate interacting Dirac fermions, one is led naturally to the problem of simulating quantum gauge field theories. Finding an algorithm with which one could simulate a quantum field theory of interacting gauge fields and fermions on a quantum computer with exponential performance enhancement would be a very interesting result. The QLGA approach is naturally suited to describing second-quantized systems in a Hamiltonian framework, so such a model is probably best discussed in the context of Hamiltonian lattice methods [24].

There are several difficulties inherent in constructing a QLGA model of a quantum gauge theory. First, it is necessary to find a lattice rule which describes the propagation of a single particle according to Maxwell’s equations. This is a challenge, because it is difficult to have an isotropic propagator on a lattice when the scaling of the spatial and temporal lattice spacing is the same. Bialynicki-Birula has suggested that his model for the Weyl spinor can be used to describe a propagating photon by writing the Maxwell equations in spinor notation. It is unclear how to formulate this model in QLGA language, however it does not seem that there is any fundamental obstacle to such a formulation. Another general difficulty with simulating gauge theories is the discretization of the gauge group. In
the QLGA formalism, however, where we are essentially dealing with a second-quantized field theory, this should not be a problem; the fermions will carry a gauge index, and the gauge particles will carry an index in the adjoint representation of the gauge group. Thus, for example, to simulate the theory of QCD with an SU(3) gauge theory, we would include three colors of fermions and 8 species of gluons, just as in the physical particle theory we are simulating.

To end this section, we briefly discuss a completely different approach to simulating quantum field theories on a quantum computer which does not rely on the QLGA framework. It is always possible to describe a field theory with a momentum cutoff by writing the Hamiltonian in terms of creation and annihilation operators for the modes of the fields with momentum below the cutoff. In a discrete time formulation, the evolution of the state of the system in a single time step is given by the exponential of the Hamiltonian times the time step (times \( i \)). If this time-development operator can be reasonably approximated by a polynomial number of two-particle interactions, then such a system could be simulated on a quantum computer with an exponential performance increase using the methods discussed in [4]. Whether such a nonlocal approach to the simulation is preferable to the local QLGA method is not clear; further study of both approaches will probably be needed to resolve this issue.

5 Conclusions

We have discussed algorithms for simulating quantum systems on quantum computers. In particular, we have described in detail algorithms which enable the simulation of interacting many-body quantum systems with exponential performance enhancement over classical computers. There are many obstacles to building the sort of large-scale quantum computers which would be necessary for the algorithms discussed here to be useful. In addition to the technical difficulties involved with precisely controlling systems of many quantum elements, there are the theoretical problems of making algorithms robust against imprecision and loss
of coherence which must appear in any physical system. Many of the other speakers at this conference have discussed these and related issues in great detail, so we have concentrated here on describing algorithms under the assumption that eventually the technical problems will be solved and we will have a programmable universal quantum computer at our disposal. Certainly, recent progress in quantum error correcting codes [25] gives some indication that there is room for optimism in this direction. Even if a true “universal quantum computer” can never be built, however, it may be possible to construct quantum devices which have some computational capacity [11]. Because the algorithms we have discussed here are based only on simple local quantum interactions, it seems plausible that many different types of quantum systems may be useful in simulating other quantum systems in a precise and relatively controlled fashion.

One interesting aspect of algorithms which simulate quantum systems on a quantum computer is the issue of measurement. In a classical computation, all the bits in the system are available at all times to the programmer, so that the state of the system can be completely known. In a quantum computer, on the other hand, the state of the system can only be known through the measurement of certain q-bits, whose values are determined probabilistically through the usual rules of quantum mechanics. This means that some information in the state of a quantum computer is not directly accessible to the programmer. In terms of the algorithms discussed here for simulating quantum systems, this means that only certain questions can be asked of the simulation. For example, imagine simulating a system of a single electron in an external potential well. In principle, given the exact time-development of the wavefunction in the quantum computer doing the simulation, a spectral analysis would immediately enable one to calculate the entire energy spectrum of the system. However, this complete information is not available. Instead, we can only ask the types of questions which we could ask in the real physical system—for example, we could ask for the probability that a particle which starts at a fixed point in space moves to a given region in a fixed time. This limitation on the set of questions which can be meaningfully asked of a quantum computer may seem at first to make the process of extracting meaningful results from such simulations
more difficult. However, since the questions which can be asked of the simulation are precisely those questions which can be asked of the system being simulated, the connection to results of physical experiments is very direct. Any difficulty in connecting the results of the simulation with predictions about physical systems must be the consequence of theoretical prejudices.

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