Implementation of a quantum algorithm on a nuclear magnetic resonance quantum computer

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Quantum computing shows great promise for the solution of many difficult problems, such as the simulation of quantum systems and the factorization of large numbers. While the theory of quantum computing is fairly well understood, it has proved difficult to implement quantum computers in real physical systems. It has recently been shown that nuclear magnetic resonance (NMR) can be used to implement small quantum computers using the spin states of nuclei in carefully chosen small molecules. Here we demonstrate the use of a NMR quantum computer based on the pyrimidine base cytosine, and the implementation of a quantum algorithm to solve Deutsch’s problem (distinguishing between constant and balanced functions). This is the first successful implementation of a quantum algorithm on any physical system. © 1998 American Institute of Physics.

I. INTRODUCTION

In 1982 Feynman pointed out that it appears to be impossible to efficiently simulate the behavior of a quantum mechanical system with a computer. This problem arises because the quantum system is not confined to its eigenstates, but can exist in any superposition of them, and so the space needed to describe the system is very large. To take a simple example, a system comprising $N$ two-level subsystems, such as $N$ spin-$\frac{1}{2}$ particles, inhabits a Hilbert space of dimension $2^N$, and evolves under a series of transformations described by matrices containing $4^N$ elements. For this reason it is impractical to simulate the behavior of spin systems containing more than about a dozen spins.

The difficulty of simulating quantum systems using classical computers suggests that quantum systems have an information processing capability much greater than that of corresponding classical systems. Thus, it might be possible to build quantum mechanical computers, which utilize this information processing capability in an effective way to achieve a computing power well beyond that of a classical computer. Such a quantum computer could be used to efficiently simulate other quantum mechanical systems or to solve conventional mathematical problems, which suffer from a similar exponential growth in complexity, such as factoring. Experimental implementation of a quantum computer has, however, proved difficult. Much effort has been directed toward implementing quantum computers using ions trapped by electric and magnetic fields, and while this approach has shown some success, it has proved difficult to progress beyond computers containing a single two-level system (corresponding to a single quantum bit, or qubit).

Recently two separate approaches have been described for the implementation of a quantum computer using nuclear magnetic resonance (NMR). These approaches show great promise, as it has proved relatively simple to investigate quantum systems containing two or three qubits. Here we describe our implementation of a simple quantum algorithm for solving Deutsch’s problem on a two qubit NMR quantum computer.

II. QUANTUM COMPUTERS

All current implementations of quantum computers are built up from a small number of basic elements. The first of these is the qubit, which plays the same role as that of the bit in a classical computer. A classical bit can be in one of two states, 0 or 1, and similarly a qubit can be represented by any two-level system with eigenstates labeled $|0\rangle$ and $|1\rangle$. One obvious implementation is to use the two Zeeman levels of a spin-$\frac{1}{2}$ particle in a magnetic field, and we shall assume this implementation throughout the rest of this paper. Unlike a bit, however, a qubit is not confined to these two eigenstates, but can, in general, exist in some superposition of the two states. It is this ability to exist in superpositions that makes quantum systems so difficult to simulate and that gives quantum computers their power.

The second requirement is a set of logic gates, corresponding to gates such as AND, OR, and NOT in conventional
Quantum gates differ from their classical counterparts in one very important way: they must be reversible. This is because the evolution of any quantum system can be described by a series of unitary transformations, which are themselves reversible. This need for reversibility has many consequences for the design of quantum gates. Clearly, for a gate to be reversible it must be possible to reconstruct the input bits knowing only the design of the gate and the output bits, and so every input bit must be in some sense preserved in the outputs. One trivial consequence of this is that the gate must have exactly as many outputs as inputs. For this reason it is obvious that gates such as AND and OR are not reversible. It is, however, possible to construct reversible equivalents of AND and OR, in which the input bits are preserved.

Just as it can be shown that one or more gates (such as the NAND gate) are universal for classical computing (that is, any classical gate can be constructed using only wires and NAND gates), it can be shown that certain gates or combinations of gates are universal for quantum computing. In particular, it can be shown that the combination of a general single qubit rotation with the two bit "controlled-NOT" gate (CNOT) is universal. Furthermore, it is possible to build a reversible equivalent of the NAND gate, and thus to implement any classical logic operation using reversible logic.

Single qubit rotations are easily implemented in NMR, as they correspond to rotations within the subspace corresponding to a single spin, and such rotations can be achieved using radiofrequency (rf) fields. One particularly important single bit gate is the Hadamard gate, which performs the rotational transformation,

\[ |0\rangle \rightarrow \frac{|0\rangle + |1\rangle}{\sqrt{2}}, \]
\[ |1\rangle \rightarrow \frac{|0\rangle - |1\rangle}{\sqrt{2}}. \]  

The Hadamard operator can thus be used to convert eigenstates into superpositions of states. Similarly, as the Hadamard is self-inverse, it can be used to convert superpositions of states back into eigenstates for later analysis.

Two-bit gates correspond to rotations within subspaces corresponding to two spins, and thus require some kind of spin–spin interaction for their implementation. In NMR the scalar spin–spin coupling (J coupling) has the correct form, and is ideally suited for the construction of controlled gates, such as CNOT. This gate operates to invert the value of one qubit when another qubit (the control qubit) has some specified value, usually |1⟩; its truth table is shown in Table I.

Finally, it is necessary to have some way of reading out information about the final quantum state of the system, and thus obtaining the result of the calculation. In most implementations of quantum computers, this process is equivalent to determining which of two eigenstates a two-level system is in, but this is not a practical approach in NMR. It is, however, possible to obtain equivalent information by exciting the spin system and observing the resulting NMR spectrum. Different qubits correspond to different spins, and thus give rise to signals at different resonance frequencies, while the eigenstate of a spin before the excitation can be determined from the relative phase (absorption or emission) of the NMR signals.

### III. THE DEUTSCH ALGORITHM

Deutsch’s problem in its simplest form concerns the analysis of single-bit binary functions:

\[ f(x): B \rightarrow B, \]

where \( B = \{0,1\} \) is the set of possible values for a single bit. Such functions take a single bit as input, and return a single bit as their result. Clearly there are exactly four such functions, which may be described by their truth tables, as shown in Table II. These four functions can be divided into two groups: the two "constant" functions, for which \( f(x) \) is independent of \( x \) (\( f_{00} \) and \( f_{11} \)), and the two "balanced" functions, for which \( f(x) \) is zero for one value of \( x \) and unity for the other (\( f_{01} \) and \( f_{10} \)). Given some unknown function \( f \) (known to be one of these four functions), it is possible to determine which of the four functions it is by applying \( f \) to two known inputs: 0 and 1. This procedure also provides enough information to determine whether the function is constant or balanced. However, knowing whether the function is constant or balanced corresponds to only one bit of information, and so it might be possible to answer this question using only one evaluation of the function \( f \). Equivalently, it might be possible to determine the value of \( f(0) \oplus f(1) \) using only one evaluation of \( f \). (The symbol \( \oplus \) indicates addition modulo 2, and for two one bit numbers, \( a \) and \( b \), \( a \oplus b \) equals 0 if \( a \) and \( b \) are the same, and 1 if they are different.) In fact, this can be achieved as long as the calculation is performed using a quantum computer rather than a classical one.

Quantum computers of necessity use reversible logic, and so it is not possible to implement the binary function \( f \) directly. It is, however, possible to design a propagator, \( U_f \), which captures \( f \) within a reversible transformation by using a system with two input qubits and two output qubits as follows:

<table>
<thead>
<tr>
<th>( x )</th>
<th>( f_{00}(x) )</th>
<th>( f_{01}(x) )</th>
<th>( f_{10}(x) )</th>
<th>( f_{11}(x) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

### TABLE I. The truth table for the CNOT gate. The first qubit (the control qubit) is unchanged by the gate, while the second qubit is flipped if the control qubit is in state 1, effectively implementing an XOR gate.

### TABLE II. The four possible binary functions mapping one bit to another.
The two input bits are preserved \( |x\rangle\) is preserved directly, while \( y \) is preserved by combining it with \( f(x) \), the desired result, and so \( U_f \) corresponds to a reversible transformation. Note that for any one bit number \( a \), \( 0 \oplus a = a \), and so values of \( f(x) \) can be determined by setting the second input bit to 0. Using this propagator and appropriate input states, it is possible to evaluate \( f(0) \) and \( f(1) \) using

\[
|0\rangle|0\rangle \rightarrow |0\rangle|f(0)\rangle
\]

and

\[
|1\rangle|0\rangle \rightarrow |1\rangle|f(1)\rangle.
\]

(4)

(5)

The approach outlined above, in which the state of a quantum computer is described explicitly, swiftly becomes unwieldy, and it is useful to use more compact notations. One particularly simple approach is to use quantum circuits, which may be drawn by analogy with classical electronic circuits. In this approach lines are used to represent "wires" down which qubits "flow," while boxes represent quantum gates that perform appropriate unitary transformations. For example, the analysis of \( f \) can be summarised by the circuit shown in Fig. 1.

So far, this is simply using a quantum computer to simulate a classical computer implementing classical algorithms. With a quantum computer, however, it is not necessary to start with the system in some eigenstate; instead, it is possible to begin with a superposition of states. Suppose the calculation begins with the second qubit in the superposition \( (|0\rangle - |1\rangle)/\sqrt{2} \). Then

\[
|x\rangle \left(\frac{|0\rangle - |1\rangle}{\sqrt{2}}\right) U_f \left(\frac{|0\oplus f(x)\rangle - |1\oplus f(x)\rangle}{\sqrt{2}}\right),
\]

(3)

\[
= \begin{cases} 
|x\rangle \frac{|0\rangle - |1\rangle}{\sqrt{2}}, & \text{if } f(x) = 0, \\
|x\rangle \frac{|1\rangle - |0\rangle}{\sqrt{2}}, & \text{if } f(x) = 1,
\end{cases}
\]

(6)

\[
=(-1)^{f(x)}|x\rangle \frac{|0\rangle - |1\rangle}{\sqrt{2}}.
\]

(We have used the fact that \( 0 \oplus a = a \), as before, while \( 1 \oplus a = 1 \) if \( a = 0 \) and \( 0 \) if \( a = 1 \).) The value of \( f(x) \) is now encoded in the overall phase of the result, with the qubits left otherwise unchanged. While this is not particularly useful, suppose the calculation begins with the first qubit also in a superposition of states, namely \( (|0\rangle + |1\rangle)/\sqrt{2} \). Then

\[
|\psi_{(0)}\rangle = |0\rangle|1\rangle.
\]

(8)

but this description is not really appropriate. Unlike other implementations, a NMR quantum computer comprises not just a single set of spins but rather an ensemble of spins in a statistical mixture of states. Such a system is most conveniently treated using a density matrix, which can describe either a mixture or a pure state; for example,
\[
\rho_{01} = |\phi_{01}\rangle \langle \phi_{01}| = \begin{pmatrix}
0 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0
\end{pmatrix}.
\]

This density matrix can be decomposed in the product operator basis as \(\rho_{01} = (I_z - S_z - 2I_xS_x + 1/2 E)/2\). Ignoring multiples of the unit matrix (which give rise to no observable effects in any NMR experiment), this can be reached from the thermal equilibrium density matrix \((I_z + S_z)\) by a series of rf and field gradient pulses.\(^{11}\)

The unitary transformation matrix corresponding to the Hadamard operator on a single spin can be written as

\[
H = \frac{1}{\sqrt{2}} \begin{pmatrix}
1 & 1 \\
1 & -1
\end{pmatrix}.
\]

This corresponds to a 180° rotation around an axis tilted at 45° between the z and x axes. Such a rotation can be achieved directly using an off resonance pulse,\(^{13}\) or using a three pulse sandwich\(^{13}\) such as 45° - 180° - 45°. Even more simply, the Hadamard can be approximated by a 90° pulse. While this is clearly not a true Hadamard operator (for example, it is not self-inverse), its behavior is similar, and it can be used in some cases: for example, it is possible to replace the first pair of Hadamard gates in the circuit for the Deutsch Algorithm (Fig. 2) by 90° pulses and the second pair of gates by 90° pulses. Clearly, it is possible to apply the Hadamard operator either to just one of the two spins (using selective soft rf pulses\(^{20}\)) or to both spins simultaneously (using nonselective hard pulses).

The unitary transformations corresponding to the four possible propagators \(U_f\) are also easily derived. Each propagator corresponds to flipping the state of the second qubit under certain conditions as follows: \(U_{00}\), never flip the second qubit; \(U_{01}\), flip the second qubit when the first qubit is in state one; \(U_{10}\), flip the second qubit when the first qubit is in state zero; \(U_{11}\), always flip the second qubit. The first and last cases are particularly simple, as \(U_{00}\) corresponds to doing nothing (the identity operation), while \(U_{11}\) corresponds to inverting the second spin (a conventional NOT gate, or, equivalently, a 180° pulse). The second and third propagators correspond to controlled-NOT gates, which can be implemented using spin-spin couplings. For example, \(U_{01}\) is described by the matrix

\[
U_{01} = \begin{pmatrix}
1 & 0 & 0 & 0 \\
0 & 1 & 0 & 0 \\
0 & 0 & 0 & 1 \\
0 & 0 & 1 & 0
\end{pmatrix},
\]

which can be achieved using the pulse sequence

\[90S_y - \text{couple} - 90I_z - 90S_z - 90S_y,\]

where 90° indicates a 90° pulse on the second spin, couple indicates evolution under the scalar coupling Hamiltonian, \(\pi J_{IS}2I_zS_z\), for a time 1/2\(J_{IS}\), and 90I\(_z\) and 90S\(_z\) indicate either periods of free precession under Zeeman Hamiltonians or the application of composite z pulses.\(^ {20,21}\) Similarly, \(U_{10}\) can be achieved using the pulse sequence

\[90S_y - \text{couple} - 90I_z - 90S_z - 90S_y,\]

The pulse sequences described above can be implemented in many different ways, as different composite z pulses can be used, the order of some of the pulses can be varied, and in some cases different pulses can be combined together. We chose to use the implementation

\[90S_y - \frac{1}{4}J_{IS} - 180_x - \frac{1}{4}J_{IS} - 180_x - 90I_y - 90I_x - 90S_y - 90S_{zx},\]

where pulses not marked as either I or S were applied to both nuclei. The phase of the final pulse distinguishes \(U_{01}\) (for which the final pulse is \(S_{zx}\)) from \(U_{10}\) (for which it is \(S_{zx}\)).

Finally, it is necessary to consider an analysis of the final state, which could, in general, be one of the four states \(|\phi_{00}\rangle\), \(|\phi_{01}\rangle\), \(|\phi_{10}\rangle\), or \(|\phi_{11}\rangle\). In order to distinguish these states it is necessary to apply a 90° pulse and observe the NMR spectrum. The final NMR signal observed from spin \(I\) is \(I_z\), if the spin is in state 0, and \(-I_x\) if it is in state 1. For a computer implementing the Deutsch algorithm the final detection 90° pulses cancel out the two final pseudo-Hadamard 90° pulses, and thus all four pulses can be omitted (Fig. 3). The final NMR signal observed is either 1/2\(I_z\) - 1/2\(I_x\) (corresponding to \(|\phi_{00}\rangle\)) or -1/2\(I_z\) - 1/2\(I_x\) (corresponding to \(|\phi_{11}\rangle\)). Hence, it is simple to determine the value of \(f(0) \oplus f(1)\) (that is, determine whether the function is constant or balanced) by determining the relative phase of the signals from the two spins.

![Diagram](https://example.com/diagram.png)
splitting in each pair arises from the scalar coupling $J_{IS}$.

The left-hand pair of signals corresponds to the first spin ($I$), while the pair on the right-hand side correspond to the second spin ($S$). Note that the signals from both spins (which are in state $|0\rangle$, the ground state) are in absorption.

The right-hand pair of signals is in absorption, while $f(0) = 0$ for this function. (c) The result of applying $U_{f0}$; the signals from spin $S$ are now in emission, since $f(0) = 1$ for this function. (d) The result of applying $U_{f1}$; the signals from spin $S$ are once again in emission as expected.

V. EXPERIMENT

In order to demonstrate the results described above, we have constructed a NMR quantum computer capable of implementing the Deutsch algorithm. For our two-spin system we chose to use a 50 mM solution of the pyrimidine base cytosine in D$_2$O; a rapid exchange of the two amine protons and the single amide proton with the deuterated solvent leaves two remaining protons forming an isolated two-spin system. All NMR experiments were conducted at 20 °C and $pH = 7$ on a home-built NMR spectrometer at the Oxford Centre for Molecular Sciences, with a $^1$H operating frequency of 500 MHz. The observed J coupling between the two protons was 7.2 Hz, while the difference in resonance frequencies was 763 Hz. Selective excitation was achieved using Gaussian soft pulses incorporating a phase ramp to allow excitation away from the transmitter frequency. During a selective pulse the other (unexcited) spin continues to experience the main Zeeman interaction, resulting in a rotation around the $z$ axis, but the length of the selective pulses can be chosen such that the net rotation experienced by the other spin is zero. The residual HOD resonance was suppressed by low-power saturation during the relaxation delays.

This system can be used both for the implementation of classical algorithms to analyze $f(0)$ and $f(1)$ and for the implementation of the Deutsch algorithm; as shown in Fig. 3 the pulse sequences differ only in the placement of the 90° pulses. The results for the classical algorithm to determine $f(0)$ are shown in Fig. 4. The left-hand pair of signals corresponds to the first spin ($I$), while the pair on the right-hand side correspond to the second spin ($S$); the (barely visible) splitting in each pair arises from the scalar coupling $J_{IS}$. In this experiment the value of $f(0)$ is determined by setting both spins $I$ and $S$ into state $|0\rangle$, performing the calculation, and then measuring the final state of spin $S$; spin $I$ should not be affected, and so should remain in state $|0\rangle$. The phase of the reference spectrum (a) was adjusted so that signals from spin $I$ appear in absorption, and the same phase correction was applied to the other three spectra. The state of a spin after a calculation can then be determined by determining whether the corresponding signals in the spectrum are in absorption (state $|0\rangle$) or emission (state $|1\rangle$). As expected, spin $I$ does indeed remain in state $|0\rangle$, while the value of $f(0)$ (determined from spin $S$) is 0 for $U_{f00}$ and $U_{f01}$, but 1 for $U_{f10}$ and $U_{f11}$.

Clearly, our NMR quantum computer is capable of implementing this classical algorithm, as it is simple to determine $f(0)$. The other value, $f(1)$, can be determined in a very similar way (see Fig. 5). In this case spin $I$ remains in state $|1\rangle$, while $f(1)$ equals 0 for $U_{f00}$ and $U_{f10}$ and equals 1 for $U_{f01}$ and $U_{f11}$. There are, however, several imperfections visible in the results.

First, the signals are not perfectly phased: rather than exhibiting pure absorption or pure emission lineshapes, the signals have more complex shapes, including dispersive components. These arise from the difficulty of implementing perfect selective pulses, which effect the desired rotation at one spin while leaving the other spin entirely unaffected. Similarly, the selective pulses will not perfectly suppress J couplings during the excitation, leading to the appearance of antiphase contributions to the lineshape. Any practical selective pulse will be imperfect, and so will result in systematic distortions in the final result. Note that these distortions are most severe in cases (b) and (c), where the propagator is complex, containing a large number of selective pulses. Interestingly, the distortions are also more severe for the measurement of $f(0)$ (Fig. 4) than for $f(1)$ (Fig. 5); there is no simple explanation for this effect, which is due to the complex interplay of many selective pulses. We are currently seeking ways to minimize these effects.

Second, the signal intensities vary in different cases; as before, the signal loss is most severe in cases (b) and (c), corresponding to complex propagators. This is in part a consequence of imperfect selective pulses, as discussed above, but may also indicate the effects of spin relaxation, that is, decoherence of the states involved in the calculation. Decoherence is a fundamental problem, and may ultimately limit

FIG. 4. Experimental implementation of an algorithm to determine $f(0)$ on a NMR quantum computer. (a) The result of applying $U_{f00}$; as this propagator is the identity matrix this spectrum can also serve as a reference. The left-hand pair of signals corresponds to the first spin ($I$), while the pair on the right-hand side correspond to the second spin ($S$). Note that the signals from both spins (which are in state $|0\rangle$, the ground state) are in absorption. (b) The result of applying $U_{f0}$; both sets of signals are still in absorption, as $f(0) = 0$ for this function. (c) The result of applying $U_{f10}$; the signals from spin $S$ are now in emission, since $f(0) = 1$ for this function. (d) The result of applying $U_{f1}$; the signals from spin $S$ are once again in emission as expected.

FIG. 5. Experimental implementation of an algorithm to determine $f(1)$ on a NMR quantum computer; in this case the algorithm starts with spin $I$ in the excited state, $|1\rangle$, and so signals from spin $I$ are in emission. For details of the labeling see Fig. 4.
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20 R. Freeman, Spin Choreography (Spectrum, Oxford, 1997).
31 Since initial submission of this manuscript there has been considerable progress in this field, including another implementation of an algorithm to solve Deutsch’s problem, and two implementations of Grover’s quantum search algorithm.