Quantum Computing by Two-Dimensional NMR using Spin- and Transition-Selective Pulses

T.S. Mahesh †, Kavita Dorai †, Arvind ♯, and Anil Kumar †‡∗

† Department of Physics, ‡ Sophisticated Instruments Facility
Indian Institute of Science, Bangalore 560012 India

Quantum computing using two-dimensional NMR has recently been described using scalar coupling evolution technique [J. Chem. Phys., 109, 10603 (1998)]. In the present paper, we describe two-dimensional NMR quantum computing with the help of selective pulses. A number of logic gates are implemented using two and three qubits with one extra observer spin. Some many-in-one gates (or Portmanteau gates) are implemented. Toffoli gate (or AND/NAND gate) and OR/NOR gates are implemented on three qubits. Deutsch-Jozsa quantum algorithm for one and two qubits, using one extra work qubit, has also been implemented using selective pulses after creating a coherent superposition state, in the two-dimensional methodology.

Key Words: Two-dimensional NMR; quantum computation; logic gate; Toffoli gate; Deutsch Jozsa algorithm.

INTRODUCTION

Quantum computing exploits the intrinsic quantum nature of physical systems and is therefore more powerful than classical computing for a certain class of problems. While some problems like factorization gain exponential speed up (1), some others like database-search gain polynomial speed up (2). A quantum computer works on two-level quantum systems known as ‘quantum bits’ or ‘qubits’. The fact that qubits can exist in a coherent superposition of basis states is exploited in a quantum computer. However, retaining such a coherent superposition for long enough time is one of the major hurdles in quantum computation. Since nuclear spins in liquids are efficiently isolated from the rest of their environment, the coherent superposition lasts for several hundreds of milliseconds. Therefore, high resolution NMR of weakly coupled nuclear spins in liquids is one of the potential candidates for realizing such a quantum computer. Nuclear spins with \( I = 1/2 \) have generally been used since spins with \( I > 1/2 \) relax much faster. Quantum computing using one-dimensional NMR has been demonstrated by various workers (3-12).

Recently, quantum computing has been demonstrated using two-dimensional NMR (13). In this method, the states of the ‘computation’ spins (or input spins) are encoded by the transitions of an extra ‘observer’ spin (Fig.1). The observer spin is first allowed to evolve for a time \( t_1 \) during which, the input qubits remain in their initial state (Fig.2). After the frequency labelling period \( t_1 \), the computation is performed on the input qubits. The observer spin is again allowed to evolve for a time \( t_2 \) and detected. The experiment starts from a mixed input state, the computation is performed simultaneously on all the input states, and one gets a mixed output state. One can also create a superposition of input qubits in the beginning of the experiment by applying a \( \pi/2 \) pulse on the input qubits and subsequently killing the coherence by a gradient pulse (13). Various steps in two-dimensional NMR namely, preparation, evolution, mixing and detection have a close correspondence with the steps in quantum computing namely, creation of initial states, labeling of the initial states, computation and reading of output states, respectively (13) (Fig.2). In this respect, two-dimensional NMR appears to be the ‘method of choice’ for quantum computing.

An advantage of the two-dimensional method is that it resolves all the input and the output states and correlates them. This correlation between input and output states in the two-dimensional experiment makes the result of the computation graphic. For example, a SWAP gate which exchanges the states of two qubits, can be implemented in NMR by selectively interchanging populations of zero-quantum levels. Since the one-dimensional NMR spectrum of homonuclear spins after the execution of SWAP gate is indistinguishable from that of an equilibrium spectrum, the one-dimensional SWAP gate was earlier demonstrated after the creation of a non-equilibrium state (12, 14). However, as will be shown here (Fig.3), and has been demonstrated earlier (13), the two-dimensional method starts from the equilibrium state and yields a spectrum characteristic of the SWAP gate.

Two-dimensional quantum computation has been carried out so far by using scalar coupling evolution tech-
EXPERIMENTAL

A. Logic Gates

A weakly coupled 3-spin system \( I_0I_1I_2 \) can be viewed as a combination of two 2-spin subsystems with the observer spin \( I_0 \) being in state 0 or 1 (Fig.1).

There are four observer spin transitions labelled as 11, 10, 01 and 00, which correspond to the states of the input qubits \( I_1I_2 \) in these transitions. During the computation (mixing) period, various transitions of the input qubits are inverted yielding various gates as listed in Table 1. The results of 24 one-to-one reversible gates for the 3-spin system having one observer and two input qubits are shown in Fig.3. The NOP gate is implemented by doing No Operation during the computation period. In this gate, each input state corresponds to same output state after the computation. Various NOT gates are implemented by inverting one or both of input qubits \( I_1I_2 \), using spin-selective \( \pi \) pulses. XOR (or control-NOT) and XNOR gates are implemented by inverting two similar transitions of the same input spin, one in each subsystem. We label XOR and XNOR gates as XOR1, XNOR1 and XOR2, XNOR2 depending on whether the result of the operation is stored on the spin \( I_1 \) or \( I_2 \) respectively. For example, XOR1 gate is implemented by inverting transitions 001-011 and 101-111 of \( I_1 \) and the result of the gate is stored on \( I_1 \) (Table 1, Fig.3). A SWAP gate can be implemented by selectively interchanging the populations of zero quantum levels of each subsystem (Table 1, Fig.3). This can be achieved by the use of a cascade of three transition-selective, non-commuting \( \pi \) pulses on reggressively connected transitions during the computation period (12, 14). It may be noted that while all the gates of ref.(13) are reproduced here, with several new gates added, the methodology used for the computation is different except for the NOP and NOT gates.

Similar to classical Boolean expressions, quantum gates can be reduced to a sequence of simple unitary operations. Although a few basic gates such as NOT, AND, OR, XOR are sufficient to carryout a given computation, other gates are also useful since they may reduce the number of pulses. Therefore 'portmanteau gates' which perform more than one operation have been implemented in one-dimensional NMR using selective pulses (12).

![Figure 1. Energy level diagram of a 3-spin system (I_0I_1I_2) showing two 2-spin subsystems corresponding to states 0 and 1 of observer spin I_0. The transitions of I_2 are labelled by the states of input qubits (I_1I_2).](image1)

![Figure 2. Pulse scheme for the two-dimensional NMR implementation of quantum computing. The close correspondence between two-dimensional NMR and quantum computing is shown. I_0 is the observer spin and I_1, I_2 are the input qubits. The gradient pulse G_z eliminates the unwanted transverse magnetization before the computation.](image2)
Figure 3. $^1$H NMR spectrum (b) of 2,3-dibromo-propionic acid (a) in CDCl$_3$ recorded on a Bruker DRX-500 spectrometer at 300K. (c) shows observer spin ($I_0$) spectra corresponding to various gates implemented using spin- and transition-selective pulses. Pulse scheme used is shown in Fig 2 and various transitions of input qubits ($I_1$, $I_2$) inverted during the computation are listed in Table 1. The spin-selective pulses were 10ms long and the transition-selective pulses were 100-300ms long. The phase of the computation pulses were cycled through ($x$, $-x$) to suppress the distortions due to pulse imperfections. All experiments were carried out in the time domain with 256 $t_1$ values and 256 complex data points along $t_2$ and with 2 scans for each $t_1$ point. Zero filling to 512×512 complex data points was done prior to the 2D Fourier transformation. All plots are shown in magnitude mode.
TABLE 1
Various logic gates with Boolean algebra, truth table and operations performed

| GATE<sup>a</sup> | IN | OUT | Transitions Inverted<sup>b</sup> | GATE<sup>a</sup> | IN | OUT | Transitions Inverted<sup>b</sup> |
|-----------------|----|-----|-------------------------------|-----------------|----|-----|-------------------------------|
| NOP | 11 | 11 | No | NOT(I<sub>1</sub>) | 11 | 01 | All transitions of I<sub>1</sub> |
| s, t→|s, t| | | | | | |
| 01 | 01 | 01 | 01 | 00 | 10 | All transitions of I<sub>1</sub> and I<sub>2</sub> |
| 00 | 00 | 00 | 00 | 10 | 00 | |
| NOT(I<sub>2</sub>) | 11 | 10 | All transitions of I<sub>2</sub> | NOT(I<sub>1</sub>, I<sub>2</sub>) | 11 | 00 | All transitions of I<sub>1</sub> and I<sub>2</sub> |
| s, t→|s, t| | | | | | |
| 10 | 11 | 111 ↔ 111 | XOR2 | 11 | 11 | 101 ↔ 100 |
| 00 | 00 | 000 ↔ 010 | | | | |
| 01 | 01 | 011 ↔ 001 | | | | |
| 00 | 00 | 010 ↔ 011 | | | | |
| XOR1 | 11 | 11 | 100 ↔ 110 | XNOR2 | 11 | 11 | 101 ↔ 100 |
| s, t→|s⊕t, t| | | | | | |
| 10 | 10 | 110 ↔ 111 | | | | |
| 00 | 00 | 010 ↔ 011 | | | | |
| 01 | 01 | 001 ↔ 011 | | | | |
| 00 | 00 | 010 ↔ 011 | | | | |
| XNOR1 | 11 | 11 | Invert ZQ: | SWAP+NOT | 11 | 00 | Invert DQ: |
| s, t→|s⊕t, t| | | | | | |
| 11 | 01 | 101 ↔ 111 | | | | |
| 10 | 11 | 100 ↔ 111 | | | | |
| 01 | 00 | 010 ↔ 011 | | | | |
| 00 | 00 | 000 ↔ 001 | | | | |
| SWAP+XOR1 | 11 | 11 | 100 ↔ 110 | SWAP+XOR2 | 11 | 11 | 100 ↔ 101 |
| s, t→|s⊕t, s| | | | | | |
| 10 | 11 | 110 ↔ 111 | | | | |
| 01 | 00 | 010 ↔ 011 | | | | |
| 00 | 00 | 000 ↔ 001 | | | | |
| SWAP+XOR | 11 | 11 | 100 ↔ 110 | SWAP+XOR2 | 11 | 11 | 100 ↔ 110 |
| s, t→|s⊕t, t| | | | | | |
| 10 | 11 | 100 ↔ 101 | | | | |
| 01 | 00 | 010 ↔ 011 | | | | |
| 00 | 00 | 000 ↔ 001 | | | | |
| SWAP+NOT+XOR | 11 | 11 | 100 ↔ 110 | SWAP+NOT+XOR2 | 11 | 11 | 100 ↔ 101 |
| s, t→|s, t| | | | | | |
| 10 | 10 | 000 ↔ 001 | | | | |
| 00 | 00 | 100 ↔ 101 | | | | |
| 01 | 00 | 010 ↔ 011 | | | | |
| 00 | 00 | 000 ↔ 001 | | | | |
| SWAP+NOT+XNOR1 | 11 | 10 | 100 ↔ 110 | SWAP+NOT+XNOR2 | 11 | 11 | 100 ↔ 101 |
| s, t→|s⊕t, t| | | | | | |
| 10 | 10 | 000 ↔ 001 | | | | |
| 00 | 00 | 100 ↔ 101 | | | | |
| 01 | 00 | 010 ↔ 011 | | | | |
| 00 | 00 | 000 ↔ 001 | | | | |
| NOT(I<sub>1</sub>) | 11 | 01 | All I<sub>1</sub> transitions and | NOT(I<sub>2</sub>) | 11 | 10 | All I<sub>2</sub> transitions and |
| s, t→|s, t| | | | | | |
| 10 | 00 | 111 ↔ 110 | | | | |
| 00 | 11 | 011 ↔ 010 | | | | |
| 01 | 11 | 010 ↔ 011 | | | | |
| 00 | 00 | 001 ↔ 000 | | | | |
| NOT(I<sub>2</sub>) | 11 | 00 | All I<sub>2</sub> transitions and | NOT(I<sub>1</sub>, I<sub>2</sub>) | 11 | 10 | All I<sub>2</sub> transitions and |
| s, t→|s⊕t, t| | | | | | |
| 10 | 00 | 111 ↔ 110 | | | | |
| 00 | 11 | 011 ↔ 010 | | | | |
| 01 | 11 | 010 ↔ 011 | | | | |
| 00 | 00 | 001 ↔ 000 | | | | |
| SWAP+NOT | 11 | 01 | Invert | SWAP+NOT | 11 | 10 | Invert |
| s, t→|t, s| | | | | | |
| 10 | 01 | | | | | |
| 01 | 00 | | | | | |
| 00 | 00 | | | | | |

<sup>a</sup> s, t represents the state of input qubits (I<sub>1</sub>, I<sub>2</sub>).
<sup>b</sup> Order of transitions are important as all transitions may not commute.

Similar gates have been implemented here using two-dimensional NMR (Table 1, Fig.3). Interchanging populations of double quantum levels using a cascade of non-commuting π pulses on progressively connected transitions leads to SWAP+NOT gate (Table 1, Fig.3). SWAP+XOR and SWAP+XNOR gates have been implemented by inverting two pairs of progressively connected transitions in each case (Table 1, Fig.3). Similarly, SWAP+NOT+XOR and SWAP+NOT+XNOR gates have been implemented by inverting two pairs of progressively connected transitions (Table 1, Fig.3). The last six gates in Fig.3 are direct combinations of two gates. It may be noted that the 24 gates shown in Fig.3 form a complete set of 2-qubit one-to-one mappings.
Fig. 4 shows several 3-qubit gates implemented on a 4-spin system, using selective pulses. Once again, No OPeration during computation period yields NOP gate and inverting spin $I_1$ yields the NOT($I_1$) gate. The more interesting ones are - Toffoli gate (or AND/NAND gate) and OR/NOR gate. The operations of Toffoli and OR/NOR gates (13, 15) are respectively,

$$|s, t, u\rangle \rightarrow |s \oplus (t \land u), t, u\rangle \quad (1)$$

and

$$|s, t, u\rangle \rightarrow |s \oplus (t \lor u), t, u\rangle, \quad (2)$$

where $\oplus \equiv$ addition modulo 2, $\land \equiv$ AND, $\lor \equiv$ OR and $s, t, u$ are the states of the control spin $I_1$ and the input spins $I_2$ and $I_3$. The Toffoli gate is a universal gate for reversible computation. This gate can be implemented by inverting the control spin transitions 011 and 111, by using a pair of transition-selective $\pi$ pulses. Inverting all transitions of the control spin except the transitions 010 and 110 leads to OR/NOR gate (Fig.4).

**B. Deutsch-Jozsa algorithm**

The Deutsch-Jozsa (DJ) algorithm demonstrates the power of quantum computing over classical computing (16). In this algorithm, we consider functions from N-bit domain space to 1-bit range space. A function is called constant if it gives the same output for any input, and is called balanced if it gives one output for half the number of inputs and another for the remaining half. Given an N-bit binary function which is either constant or balanced, the single-bit DJ algorithm are listed in Table 2. The four possible functions for the single-bit DJ algorithm is encoded in a unitary transformation by including an extra work qubit $|r\rangle$ such that

$$|r\rangle|s\rangle \rightarrow |r\rangle f(s)|s\rangle, \quad (3)$$

where $|r\rangle$ and $|s\rangle$ are respectively the states of work qubit ($I_0$) and input qubit ($I_1$). The four possible functions for the single-bit DJ algorithm are listed in Table 2.

**TABLE 2**

The four possible binary functions $(f_1-f_4)$ for 1-bit DJ algorithm

| s   | $f_1$ | $f_2$ | $f_3$ | $f_4$ |
|-----|------|------|------|------|
| 0   | 0    | 1    | 0    | 1    |
| 1   | 0    | 1    | 1    | 0    |

**Figure 4.** $^{19}$F and $^1$H NMR spectra (b) of 2,3-difluoro-6-nitrophenol (a) in CDCl$_3$ (with one drop of D$_2$O to induce the exchange of the hydroxy proton and hence to suppress its coupling to flourine nuclei) recorded on a Bruker DRX-500 spectrometer at 300K. (c) shows $^{19}$F spectra of observer spin $I_0$ corresponding to various gates. Pulse scheme used is shown in Fig.2 and various transitions of control qubit $I_1$ inverted during the computation period are described in the text. The spin-selective pulses were 1ms long and the transition-selective pulses were 200ms long. The phase of the computation pulses were cycled through ($x, -x$) to suppress distortions due to pulse imperfections. All experiments were carried out in the time domain with 1024 $t_1$ values and 256 complex data points along $t_2$ and with 2 scans for each $t_1$ point. Zero filling to 1024×1024 complex data points was done prior to the 2D Fourier transformation. All plots are shown in magnitude mode.
This algorithm has been implemented on the thermal equilibrium state and does not require the creation of a pure initial state (10,12). The two-dimensional pulse scheme used for implementing DJ algorithm is shown in Fig.5. The experiment begins with both qubits in a superposition of states, achieved by a non-selective (π/2)$_y$ pulse. This is followed by an evolution period $t_1$, propagator $U_f$ and detection period $t_2$. The transformations corresponding to $f_1$ and $f_2$ are respectively, a unity operation and a spin selective π$_x$ pulse on the work qubit. The transformations corresponding to $f_3$ and $f_4$ are implemented by transition-selective π$_x$ pulses respectively on the 10-11 and 00-01 transitions of the work qubit (12). The results of the algorithm for all the 4 functions are shown in Fig.6. The constant or balanced nature of the function is identified by the presence or absence of signal from the input qubit (I$_1$) (Fig.6). The expected signals are also shown schematically in Fig.6. For two of the functions $f_2$ and $f_4$, the expected signals are calculated in Table 3. The spin-states ($|r\rangle|s\rangle$) before computation can be paired (connected by a curved line) in such a way that each pair represents a transition of the input qubit (I$_0$). Each transition of the input qubit is labelled by the state of the remaining spin i.e., work qubit. Corresponding output pairs of spin-states after computation can be calculated by using expression [3]. From each output pair, the corresponding transition of the input spin after computation can be identified. If two spins have flipped in the output pair, then the transition becomes non-observable and will be labelled as zero quantum (ZQ) or double quantum (DQ).

**Figure 5.** Pulse scheme for the two-dimensional NMR implementation of DJ algorithm. I$_0$ is the work qubit and I$_1$,$I_2$ are the input qubits.

**Figure 6.** $^1$H NMR spectrum (a) of 5-nitro furfuraldehyde (b) in C$_6$D$_6$ on a Bruker DRX-500 spectrometer at 300K. The results of DJ algorithm for various functions $f_1$-$f_4$ in Table 2 are shown in (c). I$_0$ is the work qubit and I$_1$ is the input qubit. Expansions of the I$_1$ parts of the spectra are shown in dotted boxes. The expected pattern is also shown in each case. The expected patterns for $f_2$ and $f_4$ are described in Table 3. The spin-selective pulses were 10ms long and the transition-selective pulses were 100ms long. The phase of the computation pulses were cycled through (x,$-x$) to suppress the distortions due to pulse imperfections. All experiments were carried out in the time domain with 256 $t_1$ values and 512 complex data points along $t_2$ and with 2 scans for each $t_1$ point. Zero filling to 512×512 complex data points was done prior to the 2D Fourier transformation. All plots are shown in magnitude mode.
TABLE 3
The input-output correlations for the functions \( f_2 \) and \( f_4 \) of 1-bit DJ algorithm.

| \( |r\rangle |s\rangle \) s-transitions | \( f_2 \) \( |f_2 \oplus r\rangle |s\rangle \) s-transitions |
|---|---|---|
| 1 1 | 0 1 | 1 0 |
| 1 0 | 1 0 | 0 1 |
| 0 1 | 1 1 | 0 1 |
| 0 0 | 1 0 | 1 1 |

TABLE 4.
The eight possible binary functions (\( f_1 - f_8 \)) for the 2-bit DJ algorithm.

| \( s \) | \( t \) | \( f_1 \) | \( f_2 \) | \( f_3 \) | \( f_4 \) | \( f_5 \) | \( f_6 \) | \( f_7 \) | \( f_8 \) |
|---|---|---|---|---|---|---|---|---|---|
| 0 0 | 0 0 | 1 0 1 1 0 1 0 1 |
| 0 1 | 0 1 0 1 1 0 1 1 |
| 1 0 | 0 1 1 0 0 0 1 1 |
| 1 1 | 0 1 0 0 0 0 0 0 |

Implementing DJ-algorithm on two input qubits requires three qubits including one work qubit. The algorithm can be described as

\[
|r\rangle |s\rangle |t\rangle \xrightarrow{U_f} |r \oplus f(s, t)\rangle |s\rangle |t\rangle,
\]

where \( |r\rangle, |s\rangle \) and \( |t\rangle \) are the states of the work qubit (I_0) and of two input (I_1, I_2) qubits respectively. The eight possible two-bit binary functions are listed in Table 4.

Figure 7. The results of DJ algorithm on 2,3-dibromo-propionic acid (Fig.3a,b) in CDCl_3 for various functions \( f_1 - f_8 \) listed in Table 4. I_0 is the work qubit and I_1, I_2 are the input qubits. Only expansions of the I_1 and I_2 parts of the spectra are shown. Expected pattern is also shown in each case for comparison. The expected patterns for \( f_2 \) and \( f_4 \) are described in Table 5. All experiments were carried out on a Bruker DRX-500 spectrometer at 300K. The spin-selective pulses were 10ms long and the transition-selective pulses were 100-300ms long. The phase of the computation pulses were cycled through \((x, -x)\) to suppress the distortions due to pulse imperfections. 1024 complex data points along \( t_2 \) were acquired for 256 \( t_1 \) values with 2 scans for each \( t_1 \) point. Zero filling to 1024×1024 complex data points was done prior to the 2D Fourier transformation. All plots are shown in magnitude mode.

Once again, a function is constant only if signals from all the input qubits are present, otherwise the function is balanced (Fig.7). Table 5 describes the input-output correlation for \( f_2 \) and \( f_4 \) in a 2-qubit DJ algorithm. Here the s- and t- transitions are labelled by the states \( |r\rangle |t\rangle \) and \( |r\rangle |s\rangle \) respectively. Other details of Table 5 are similar to that of Table 3.
The use of heteronuclear spins such as the selective pulse experiments, one needs resolved transitions in order to achieve selectivity may be difficult to achieve. Furthermore, for 2-qubit DJ algorithm on 2 and 3 weakly coupled spin systems by utilizing spin- and transition-selective pulses. The use of selective pulses over scalar evolution has several advantages as well as some disadvantages. The advantages of selective pulses are that they lead to simple logic, need only a few pulses and work very well. The disadvantages are that, one needs long low power r.f. pulses during which relaxation and r.f. inhomogeneity effects degrade the performance of the selective pulses and ideal selectivity may be difficult to achieve. Furthermore, for the selective pulse experiments, one needs resolved transitions. However, many of these difficulties can be overcome by the use of heteronuclear spins such as $^{15}$N, $^{13}$C, $^{19}$F, $^{31}$P and $^1$H, where the magnitudes of the spin-spin couplings are large, the transitions are well spread out and selectivity is easy to achieve using pulses of shorter durations. Further improvement can be achieved by orienting molecules in liquid crystal matrices. Attempts are continuing in this direction in our as well as in other laboratories (18,19).

**CONCLUSIONS**

Using two-dimensional NMR quantum computing method, several 2- and 3-qubit gates have been implemented on 3 and 4 weakly coupled spin systems and 1- and 2-qubit DJ algorithm on 2 and 3 weakly coupled spin systems by utilizing spin- and transition-selective pulses. The use of selective pulses over scalar evolution has several advantages as well as some disadvantages. The advantages of selective pulses are that they lead to simple logic, need only a few pulses and work very well. The disadvantages are that, one needs long low power r.f. pulses during which relaxation and r.f. inhomogeneity effects degrade the performance of the selective pulses and ideal selectivity may be difficult to achieve. Furthermore, for the selective pulse experiments, one needs resolved transitions. However, many of these difficulties can be overcome by the use of heteronuclear spins such as $^{15}$N, $^{13}$C, $^{19}$F, $^{31}$P and $^1$H, where the magnitudes of the spin-spin couplings are large, the transitions are well spread out and selectivity is easy to achieve using pulses of shorter durations. Further improvement can be achieved by orienting molecules in liquid crystal matrices. Attempts are continuing in this direction in our as well as in other laboratories (18,19).

**ACKNOWLEDGMENT**

The use of 400 and 500 MHz FTNMR spectrometers of the Sophisticated Instruments Facility (SIF), Indian Institute of Science, funded by Department of Science and Technology, New Delhi are gratefully acknowledged. We also thank Prof. K. V. Ramanathan and Dr. G. A. Naganagowda of SIF for discussions.

**REFERENCES**

1. P. W. Shor, Polynomial-time algorithms for prime factorization and discrete algorithms on quantum computer, SIAM Rev. 41, 303-332 (1999).

2. L. K. Grover, Quantum mechanics helps in searching for a needle in a haystack, Phys. Rev. Lett. 79, 325-328 (1997).

3. D. G. Cory, A. F. Fahmy, and T.F. Havel, Ensemble quantum computing by NMR spectroscopy, Proc. Natl. Acad. Sci. USA 94, 1634-1639 (1997).

4. N. Gershenfeld and I. L. Chuang, Bulk spin-resonance quantum computation, Science 275, 350-356 (1997).

5. I. L. Chuang, L. M. K. Vandersypen, X. Zhou, D. W. Leung and S. Lloyd, Experimental realization of a quantum algorithm, Nature 393, 143-146 (1998).

6. I. L. Chuang, N. Gershenfeld, and M. Kubinec, Experimental implementation of fast quantum searching, Phys. Rev. Lett. 80, 3408-3411 (1998).

7. I. L. Chuang, N. Gershenfeld, M. G. Kubinec, and D. W. Leung, Bulk quantum computation with nuclear magnetic resonance: theory and experiment, Proc. Roy. Soc. Lond. A 454, 447-467 (1998).

8. J. A. Jones, R. H. Hansen, and M. Mosca, Quantum logic gates and nuclear magnetic resonance pulse sequences, J. Magn. Reson. 135, 353-360 (1998).

9. J. A. Jones and M. Mosca, Implementation of a quantum algorithm on a nuclear magnetic resonance quantum computer, Jl. Chem. Phys. 109, 1648-1653 (1998).

10. N. Linden, H. Barjat and R. Freeman, An implementation of the Deutsch-Jozsa algorithm on a three-qubit NMR quantum computer, Chem. Phys. Lett. 296, 61-67 (1998).

11. Arvind, Kavita Dorai, and Anil Kumar, Quantum entanglement in the NMR implementation of the Deutsch-Jozsa algorithm, quant-ph/9909067.

12. Kavita Dorai, Arvind, and Anil Kumar, Implementing quantum-logic operations, pseudopure states, and the Deutsch-Jozsa algorithm using non-commuting selective pulses in NMR, Phys. Rev. A 61, 042306/1-7 (2000).
13. Z. L. Madi, R. Bruschweiler, and R. R. Ernst, One- and two-dimensional ensemble quantum computing in spin Liouville space, J. Chem. Phys. 109, 10603-10611 (1998).

14. Kavita Dorai and Anil Kumar, Cascades of selective pulses on connected single-quantum transitions leading to the selective excitation of multiple-quantum coherences, J. Magn. Reson. 114, 155-162 (1995).

15. T. Toffoli, Languages and Programming, in “Automata” (J. W. de Bakker and J. van Leeuwen, Eds.), pp. 632-644, Springer (1980).

16. D. Deutsch and R. Jozsa, Rapid solution of problems by quantum computation, Proc. Roy. Soc. Lond. A 439, 553-558 (1992).

17. R. Cleve, A. Ekert, C. Macchiavello, and M. Mosca, Quantum algorithms revisited, Proc. Roy. Soc. Lond. A 454, 339-354 (1998).

18. C. S. Yannoni, M. H. Sherwood, D. C. Miller, I. L. Chuang, L. M. K. Vandersypen, L. M. K. Vandersypen, and M. G. Kubinec, Nuclear magnetic resonance quantum computing using liquid crystal solvents, Appl. Phys. Lett. 75, 3563-3565 (1999).

19. M. Marjanska, I. L. Chuang, and M. G. Kubinec, Demonstration of quantum logic gates in liquid crystal nuclear magnetic resonance, J. Chem. Phys. 112, 5095-5099 (2000).